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Prognostic models for newly-diagnosed chronic lymphocytic leukaemia in adults: a systematic review and meta-analysis (Review)

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[Prognosis Review]

Prognostic models for newly-diagnosed chronic lymphocytic leukaemia in adults: a systematic review and meta-analysis

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ABSTRACT

Background

Chronic lymphocytic leukaemia (CLL) is the most common cancer of the lymphatic system in Western countries. Several clinical and biological factors for CLL have been identified. However, it remains unclear which of the available prognostic models combining those factors can be used in clinical practice to predict long-term outcome in people newly-diagnosed with CLL.

Objectives

To identify, describe and appraise all prognostic models developed to predict overall survival (OS), progression-free survival (PFS) or treatment-free survival (TFS) in newly-diagnosed (previously untreated) adults with CLL, and meta-analyse their predictive performances.

Search methods

We searched MEDLINE (from January 1950 to June 2019 via Ovid), Embase (from 1974 to June 2019) and registries of ongoing trials (to 5 March 2020) for development and validation studies of prognostic models for untreated adults with CLL. In addition, we screened the reference lists and citation indices of included studies.

Selection criteria

We included all prognostic models developed for CLL which predict OS, PFS, or TFS, provided they combined prognostic factors known before treatment initiation, and any studies that tested the performance of these models in individuals other than the ones included in model development (i.e. 'external model validation studies'). We included studies of adults with confirmed B-cell CLL who had not received treatment prior to the start of the study. We did not restrict the search based on study design.



Data collection and analysis

We developed a data extraction form to collect information based on the Checklist for Critical Appraisal and Data Extraction for Systematic Reviews of Prediction Modelling Studies (CHARMS). Independent pairs of review authors screened references, extracted data and assessed risk of bias according to the Prediction model Risk Of Bias ASsessment Tool (PROBAST). For models that were externally validated at least three times, we aimed to perform a quantitative meta-analysis of their predictive performance, notably their calibration (proportion of people predicted to experience the outcome who do so) and discrimination (ability to differentiate between people with and without the event) using a random-effects model. When a model categorised individuals into risk categories, we pooled outcome frequencies per risk group (low, intermediate, high and very high). We did not apply GRADE as guidance is not yet available for reviews of prognostic models.

Main results

From 52 eligible studies, we identified 12 externally validated models: six were developed for OS, one for PFS and five for TFS. In general, reporting of the studies was poor, especially predictive performance measures for calibration and discrimination; but also basic information, such as eligibility criteria and the recruitment period of participants was often missing. We rated almost all studies at high or unclear risk of bias according to PROBAST. Overall, the applicability of the models and their validation studies was low or unclear; the most common reasons were inappropriate handling of missing data and serious reporting deficiencies concerning eligibility criteria, recruitment period, observation time and prediction performance measures.

We report the results for three models predicting OS, which had available data from more than three external validation studies:

CLL International Prognostic Index (CLL-IPI)

This score includes five prognostic factors: age, clinical stage, IgHV mutational status, B2-microglobulin and TP53 status. *Calibration*: for the low-, intermediate- and high-risk groups, the pooled five-year survival per risk group from validation studies corresponded to the frequencies observed in the model development study. In the very high-risk group, predicted survival from CLL-IPI was lower than observed from external validation studies. *Discrimination*: the pooled c-statistic of seven external validation studies (3307 participants, 917 events) was 0.72 (95% confidence interval (CI) 0.67 to 0.77). The 95% prediction interval (PI) of this model for the c-statistic, which describes the expected interval for the model's discriminative ability in a new external validation study, ranged from 0.59 to 0.83.

Barcelona-Brno score

Aimed at simplifying the CLL-IPI, this score includes three prognostic factors: IgHV mutational status, del(17p) and del(11q). *Calibration*: for the low- and intermediate-risk group, the pooled survival per risk group corresponded to the frequencies observed in the model development study, although the score seems to overestimate survival for the high-risk group. *Discrimination*: the pooled c-statistic of four external validation studies (1755 participants, 416 events) was 0.64 (95% CI 0.60 to 0.67); 95% PI 0.59 to 0.68.

MDACC 2007 index score

The authors presented two versions of this model including six prognostic factors to predict OS: age, B2-microglobulin, absolute lymphocyte count, gender, clinical stage and number of nodal groups. Only one validation study was available for the more comprehensive version of the model, a formula with a nomogram, while seven studies (5127 participants, 994 events) validated the simplified version of the model, the index score. *Calibration*: for the low- and intermediate-risk groups, the pooled survival per risk group corresponded to the frequencies observed in the model development study, although the score seems to overestimate survival for the high-risk group. *Discrimination*: the pooled c-statistic of the seven external validation studies for the index score was 0.65 (95% CI 0.60 to 0.70); 95% PI 0.51 to 0.77.

Authors' conclusions

Despite the large number of published studies of prognostic models for OS, PFS or TFS for newly-diagnosed, untreated adults with CLL, only a minority of these (N = 12) have been externally validated for their respective primary outcome. Three models have undergone sufficient external validation to enable meta-analysis of the model's ability to predict survival outcomes. Lack of reporting prevented us from summarising calibration as recommended. Of the three models, the CLL-IPI shows the best discrimination, despite overestimation. However, performance of the models may change for individuals with CLL who receive improved treatment options, as the models included in this review were tested mostly on retrospective cohorts receiving a traditional treatment regimen. In conclusion, this review shows a clear need to improve the conducting and reporting of both prognostic model development and external validation studies. For prognostic models to be used as tools in clinical practice, the development of the models (and their subsequent validation studies) should adapt to include the latest therapy options to accurately predict performance. Adaptations should be timely.

PLAIN LANGUAGE SUMMARY

How well do tools predict what happens with adults with newly-diagnosed chronic lymphocytic leukaemia (CLL) over time?

What was the aim of this review?



There are many types of blood cancers called leukaemia. Chronic lymphocytic leukaemia (CLL) is the most common type. Twenty-five per cent of people who have leukaemia have CLL. It is natural for people with newly-diagnosed CLL and their families to want to know what will happen with their health in the future. They may be wondering if or when they will need treatment, if or when their disease will get worse or how long people live with CLL.

Researchers identified several characteristics that are associated with these outcomes. From these characteristics, they have tried to design tools that help predict what may happen to groups of people with newly-diagnosed CLL.

The aim of this Cochrane Review is to evaluate and summarise those tools and studies that test the tools with other patient data.

What are the key messages from this review?

Reviewers found that there is no reliable way to predict what might happen over time to people who have (untreated) CLL. One reason is because the prediction tools have not been tested enough times with enough different people to know how well they really work.

Another reason is because researchers continue to develop more effective CLL treatment options that have better results, and the prediction tools have not kept up with advances in treatment.

What are the main results of the review?

We identified 52 tools that were designed to predict what may happen to people newly-diagnosed with CLL. To find the best tools, we had to select the studies carefully. To apply these tools in clinical practice:

- a tool has to be tested by different researchers to predict what may happen with individuals with CLL in different geographic locations using different groups of people (i.e. age, gender, stage) with CLL. In other words, we would not include a tool if it was only tested on the people who provided their data to create it;
- the results of the tool should be consistent to prove that it works;
- the tests of the tool have to provide enough information to show how well the tool works. For example, the tests have to include large groups of people and enough information about the type of CLL they have.

We found three tools that met these requirements: the CLL International Prognostic Index (CLL-IPI), the Barcelona-Brno score, and the MDACC 2007 index score.

The CLL-IPI did the best job at identifying people who would survive longer with CLL and people who would survive less long. However, we rated the quality of the CLL-IPI studies as low because they did not provide all the information necessary to know how accurate the tool was. The Barcelona-Brno score and the MDACC 2007 index score, tested on a smaller overall number of patients, showed lower discrimination between persons with a good as compared to a worse prognosis, and showed a similarly low quality of the studies.

Conclusion

More and better research is needed to develop and test the tools to help predict how CLL will behave for different groups of people over time. The tools must also adapt to accurately predict the performance of new treatments.

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Summary of findings 1. CLL International Prognostic Index

Prognostic models for chronic lymphocytic leukaemia in adult patients

Population: untreated individuals with CLL **Index model:** CLL international prognostic index

Timing: moment of prediction at diagnosis of CLL; moment of outcome occurrence not prespecified (any moment after diagnosis was included)

Setting: inpatient and outpatient care

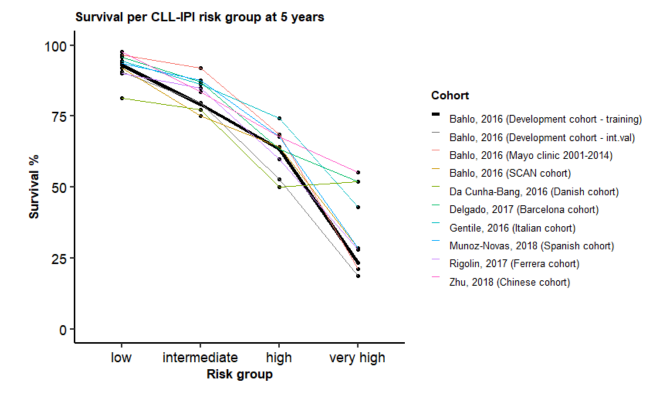
Outcomes	Measure	№ of participants (studies)	Summary mea- sure	Pooled resul	t (95% CI)			Comments
		(studies)	Juie	risk group	n (deaths)	pooled result (95% CI)	95%-PI	
Overall survival (OS) ^a	Discrimina- tion	7 studies 3307 patients 917 deaths	c-statistic of ex- ternal valida- tion studies			0.72 (95% CI 0.67 to 0.77)	0.59 to 0.83	GRADE ^c
	Calibration ^b	8 studies	survival per risk group	low	2497 (249)	92.5% (89.2% to 94.8%)	82.4% to 97.0%	Survival at 5 years
		4891 patients 875 deaths		intermedi- ate	1428 (233)	85.0% (79.7% to 89.1%)	68.8% to 93.6%	GRADE ^C
				high	765 (280)	64.9% (56.4% to 72.6%)	41.9% to 82.6%	
				very high	201 (113)	40.4% (29.3% to 52.6%)	13.2% to 72.4%	

CI: confidence interval; PI: prediction interval

^qThe CLL-IPI was developed to predict overall survival. Although we identified external validation studies for other outcomes, we limited our analysis to the primary outcome of the development study.

^bNo calibration measures were reported, and so we used data on survival frequencies per group to compare expected versus observed survival – see Figure 19. ^cGRADE was not conducted, as currently, no GRADE guidance for prognostic models exists.

Figure 19. Representation of survival per risk group per development and external validation study of the CLL-IPI (Bahlo 2016)



Summary of findings 2. Barcelona-Brno model

Prognostic models for chronic lymphocytic leukaemia in adult patients

Population: untreated individuals with CLL **Index model:** Barcelona-Brno model

Timing: moment of prediction at diagnosis of CLL; moment of outcome occurrence not prespecified (any moment after diagnosis was included)

Setting: inpatient and outpatient care

Outcomes	Measure	№ of participants (studies)	Summary mea- sure	Pooled result	: (95% CI)			Comments
		(3122123)	34.5	risk group	n (deaths)	pooled result (95% CI)	95% PI	

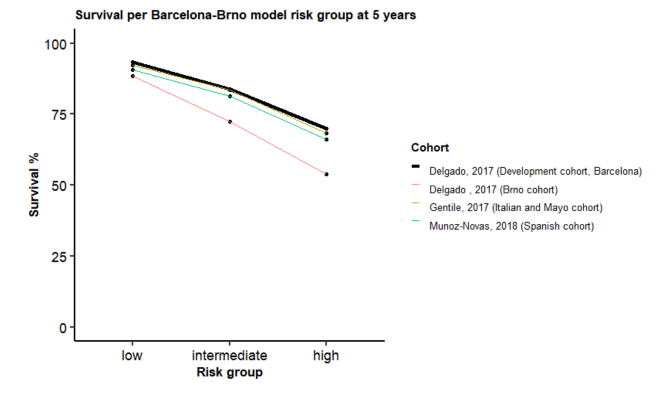
	Overall survival (OS) ^a	Discrimina- tion	4 studies 1755 patients 416 deaths	c-statistic of ex- ternal validation studies			0.64 (95% CI 0.60 to 0.67)	0.59 to 0.68	GRADE ^c
		Calibrationb	3 studies 1974 patients	survival per risk group	low	1042 (88)	90.5% (85.1% to 94.0%)	80.4% to 95.7%	Survival at 5 years
			317 deaths		intermedi- ate	673 (131)	79.7% (70.7% to 86.5%)	63.1% to 90.0%	GRADE ^c
					high	259 (98)	62.5% (49.3% to 74.1%)	41.3% to 79.7%	
٠									

CI: confidence interval; PI: prediction interval

^qThe Barcelona-Brno model was developed to predict overall survival. Although we identified external validation studies for other outcomes, we limited our analysis to the primary outcome of the development study.

^bNo calibration measures were reported, and so we used data on survival frequencies per group to compare expected versus observed survival – see Figure 20. ^cGRADE was not conducted, as currently, no GRADE guidance for prognostic models exists.

Figure 20. Representation of survival per risk group per development and external validation study of the Barcelona-Brno model (Delgado 2017)



Summary of findings 3. MDACC 2007 index score

Prognostic models for chronic lymphocytic leukaemia in adult patients

Population: untreated individuals with CLL **Index model:** MDACC 2007 index score

Timing: moment of prediction at diagnosis of CLL; moment of outcome occurrence not prespecified (any moment after diagnosis was included)

Setting: inpatient and outpatient care

Outcomes	Measure	№ of participants (studies)	sure						
		(seaules)	Juic	Risk group	n (deaths)	pooled result (95% CI)	95% PI		

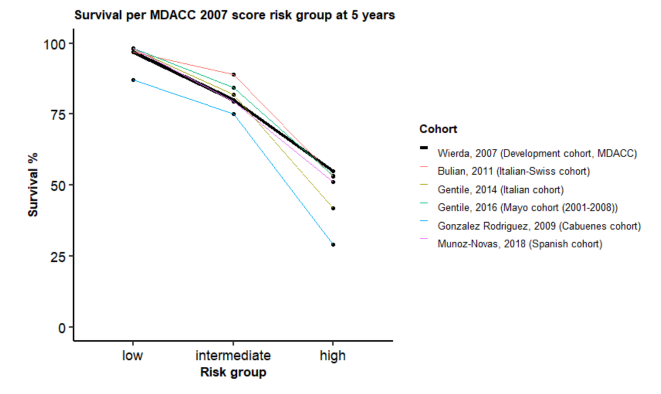
CI: confidence interval; PI: prediction interval

Overall survival (OS) ^a	Discrimina- tion	7 studies 5127 patients 994 deaths	c-statistic of ex- ternal validation studies			0.65 (95% CI 0.60 to 0.70)	0.51 to 0.77	GRADE ^c
	Calibrationb	5 studies 3786 patients	survival per risk group	low	1202 (35)	97.0% (94.3% to 98.4%)	90.9% to 99.0%	Survival at 5 years
		511 deaths		intermedi- ate	2425 (393)	82.3% (74.6% to 88.0%)	61.5% to 93.1%	GRADE ^c
				high	159 (83)	45.6% (31.3% to 60.5%)	21.2% to 72.3%	

^qThe MDACC 2007 index score was developed to predict overall survival. Although we identified external validation studies for other outcomes, we limited our analysis to the primary outcome of the development study.

^bNo calibration measures were reported, and so we used data on survival frequencies per group to compare expected versus observed survival – see Figure 21. ^cGRADE was not conducted, as currently, no GRADE guidance for prognostic models exists.

Figure 21. Representation of survival per risk group per development and external validation study of the MDACC 2007 model (Wierda 2007)







BACKGROUND

Description of the condition

Chronic lymphocytic leukaemia (CLL) is the most common form of malignant neoplasm (cancer) of the lymphatic system in Western countries. It is responsible for 25% of all leukaemias and occurs mainly in the elderly population (Chiorazzi 2005). The reported age-adjusted incidence rate of CLL in the USA between 2012 and 2016 was 4.9 per 100,000 persons with an estimated 20,720 new cases and an age-adjusted death rate of 1.2 per 100,000 persons per year (Howlader 2019). In the European Union, an estimated 46,000 individuals were living with CLL five years post-diagnosis in 2006 (Watson 2008).

In CLL, mature B cells accumulate in the blood, bone marrow, lymph nodes and spleen (Herishanu 2013). The diagnosis of CLL is generally established based on blood counts, differential counts, blood smears and immunophenotyping (Hallek 2017). The requirement for a diagnosis of CLL has been modified from a chronic absolute lymphocytosis with more than 5.0×10^9 cells per L to an absolute count of more than 5.0×10^9 monoclonal B cells with CLL immunophenotype per L peripheral blood for the duration of at least three months (Hallek 2008; Hallek 2017). In the case of a monoclonal B-cell count lower than 5.0 × 10⁹ per L and the absence of disease-related symptoms, cytopenias or tissue involvement other than bone marrow, the condition is defined as monoclonal B-lymphocytosis (Hallek 2008). A diagnosis of small lymphocytic lymphoma (SLL) is made when lymphadenopathy (enlarged lymph nodes) or splenomegaly (enlarged spleen) are caused by infiltrating CLL cells, and B lymphocytes in the peripheral blood do not exceed 5.0×10^9 per L (Hallek 2008).

CLL is characterised by a highly variable clinical course and prognosis. Some individuals with CLL experience no or only few symptoms over many years, do not require treatment, and have a life expectancy comparable to that of a healthy individual. Other individuals already experience symptoms at diagnosis or shortly thereafter, and die within a few years despite treatment. The heterogeneity in clinical presentation makes it difficult for the physician to predict accurately whether a patient may benefit from an early, aggressive treatment strategy, and to provide the patient with relevant prognostic information.

The most commonly used early attempts to group individuals with CLL according to their risk are the two staging systems by Binet 1981 (Binet 1981) and Rai 1975 (Rai 1975), which distinguish between early (Rai 0; Binet A), intermediate (Rai I, II; Binet B) and advanced stages (Rai III, IV; Binet C). The disease stage is determined by the number of lymphocytes in the peripheral blood, presence of enlarged lymph nodes, presence of anaemia or thrombocytopenia (low platelet count), and presence of an enlarged liver or spleen. The prognostic value of the two staging systems is limited as survival times vary significantly within these stages. According to the guidelines by the European Society for Medical Oncology, B-symptoms at diagnosis (fever, night sweats or weight loss) categorise people with early stage disease according to Binet or Rai, and are considered to have aggressive disease, requiring treatment (Eichhorst 2015).

Prognostic models

A prognostic model is a mathematical function that considers at least two prognostic factors, also called predictors, simultaneously with the aim to provide an estimate of an individual patient's probability to experience a certain health event within a defined time frame in the future (Alba 2017; Moons 2009; Riley 2019; Steyerberg 2013). Prognostic factors may be characteristics of the individual or the disease (e.g. age, gender, disease stage, biological or genetic information), which are likely to predict a patient-relevant outcome, such as overall survival or disease progression (Riley 2013). By statistical modelling methods, these factors are often combined to form a weighted model able to accurately predict the likelihood of this outcome. Such a model aims at assisting the clinician to estimate the patient's prognosis and enhance shared decision-making. Establishing a prognosis for the individual patient may also lead to risk-stratified treatment recommendations (Alba 2017; Debray 2017; Moons 2009; Riley 2019; Steyerberg 2013).

To develop a prognostic model for a specific outcome, such as death or disease progression, various data sources such as cohort, nested case-control or case-cohort studies are considered appropriate, especially when data are prospectively collected. Data originating from randomised clinical trials, as a special form of prospectively collected data, can be used, but may limit generalisability due to more restrictive eligibility criteria, selective participation of specialised centres, trial effects and unrealistically precise predictor assessments (Collins 2015; Moons 2014; Moons 2019; Pajouheshnia 2019). Before a prognostic model is used in clinical practice, its predictive performance should be quantified. Apparent performance of the model is the model performance estimated from the same data as used for model development and usually provides an overly optimistic estimate due to overfitting of the model to this specific dataset. Internal validation tests the model performance in the development dataset by using techniques such as bootstrapping or cross-validation. In order to test the generalisability of a prognostic model, predictive performance should be ideally assessed in several independent sets of data of individuals that were not used in the development and internal validation of the model, preferably by independent investigators to reduce bias. This process is called external validation (Moons 2015).

In the past few years, a number of clinical and biological CLL prognostic factors have been identified, including genomic aberrations, gene abnormalities (p53, ATM), mutation status of the variable segments of the immunoglobulin heavy chain genes (IgHV), or surrogate markers for these factors, such as CD38 and ZAP-70 expression (Döhner 2000; El Rouby 1993; Kay 2007). Assessing recent molecular markers at time of diagnosis is expected to provide more reliable information regarding optimal time for treatment initiation, type of therapy and individual prognosis (Kay 2007; Shanafelt 2004; Wierda 2011; Zenz 2011). Thus, progressive and smouldering forms of the disease can now be separated more accurately than by using Rai or Binet staging systems alone. Moreover, the early recognition of aggressive stage A and indolent stage B and C disease would allow rational application of risk-adapted treatment strategies. Factors influencing the choice of treatment include age, fitness to tolerate chemotherapy or immunotherapy or both, TP53 status, previous or current immune



cytopenias, and evidence of lymphomatous transformation (Goede 2012).

The recent increase in availability of biological markers for CLL presents a challenge, as well as an opportunity to develop more precise prognostic models or algorithms that integrate a combination of markers and may guide counselling and treatment decisions for the individual patient. In order to identify the best-performing tool to estimate prognosis for untreated individuals with CLL, a comprehensive evaluation of all currently available prognostic models and meta-analysis of their predictive performance in external validation studies is urgently needed.

Health outcomes

The highly variable course of CLL and the possibility of having a normal life expectancy without progression or need for treatment entails that overall survival (OS) is one of the most important outcomes to be predicted by a prognostic model. In the USA, the median age at diagnosis is 70 years and the five-year survival rate with CLL is 85.1% (Howlader 2019). Therefore, it is important to observe patients as long as possible to obtain a prognostic model that is meaningful not only for high-risk individuals, but also for people with a less aggressive disease and longer survival.

As individuals with CLL are usually older, which implies an increased prevalence of comorbidities and decreased physical fitness, treatment may lead to serious adverse events and interactions with other medications. Hence, alternative meaningful outcomes to be predicted by a prognostic model include progression-free survival (PFS) or treatment-free survival (TFS,

also sometimes referred to as time-to-first-treatment). Treatment options for CLL have improved over time, thus, affecting survival rates but not the rates of treatment indication.

Why it is important to do this review

Although several prognostic factors have been identified during the last decade (Pflug 2014; Stilgenbauer 2014; Zaja 2013), they are controversial and there is no single prognostic factor available to determine treatment options in CLL patients. These factors have also been combined into numerous prognostic models. To date, no systematic review has been conducted to evaluate and assess the predictive performance of prognostic models in CLL, which would inform us which models have the greatest validity and therefore, would be preferred to guide clinical decision-making. To shed light on this important research question, we conducted a systematic review and, where possible, a meta-analysis of existing prognostic models for CLL and their corresponding validation studies.

OBJECTIVES

To identify, describe and appraise all prognostic models developed to predict overall survival (OS), progression-free survival (PFS) or treatment-free survival (TFS) in newly-diagnosed (previously untreated) adults with CLL, and meta-analyse their predictive performances.

METHODS

Criteria for considering studies for this review

Table 1. PICOTS	system for prognostic models	
P	Population	Untreated individuals with chronic lymphocytic leukaemia (CLL) at time of prediction
1	Index model(s)	All developed prognostic models for CLL and their corresponding external validation studies
С	Comparator	No predefined comparator
0	Outcome(s)	Overall survival (OS), progression-free survival (PFS), treatment-free survival (TFS)
Т	Timing	Moment of prediction at diagnosis of CLL; moment of outcome occurrence not prespecified (any moment after diagnosis was included)
s	Setting	Not specified

Types of studies

According to the checklist for critical appraisal and data extraction for systematic reviews of prediction modelling studies (CHARMS) (Moons 2014), we included:

- prognostic model development studies without external validation in independent data;
- prognostic model development studies with external validation in independent data reported in the same study; and

 external model validation studies of previously reported models.

A prognostic model was defined as some form of mathematical function including at least two independent factors to predict OS, PFS or TFS. We included only models that combined prognostic factors known and assessed before treatment initiation. We excluded prognostic factor finding studies (i.e. studies aiming to establish one or several variables as independent prognostic factor(s) associated with an outcome, but not aiming to develop



a model to be used for individualised predictions (i.e. to provide absolute prognostic outcome probabilities) in new patients, and model impact studies (i.e. studies aiming to investigate the impact of the use of a model in practice) (Bouwmeester 2012). We excluded studies that were published only as conference proceedings. We did not exclude studies with treatment during follow-up time.

Participants

We included studies on individuals with previously untreated, confirmed B-cell CLL. We included both male and female adults (age \geq 18 years).

Types of outcomes to be predicted

Primary outcome

· Models that predict OS as an outcome

We chose this outcome because it has the greatest clinical relevance and is most important for patients. Furthermore, death due to any cause is an objective endpoint not susceptible to bias of the outcome assessor. We did not require studies to have a minimum follow-up time for inclusion in this review.

Secondary outcomes

• Models that predict either PFS or TFS

We chose these outcomes as patients with similar survival may nevertheless have differing lengths of time without symptoms or need for treatment, depending both on initial treatment and disease characteristics. In case of immediate start of treatment, identification of patients with a lower probability to obtain a good response will help in making decisions regarding treatment, for example, deciding which patients might receive new or more aggressive therapy regimens. In case of a watch-and-wait strategy, differences in estimated prognosis can influence patient management regarding surveillance and treatment.

Search methods for identification of studies

Electronic searches

Reporting and therefore retrieval of prognostic model studies is very poor, as guidelines on reporting of prediction models have only recently been published (Collins 2015). For our first search, we did not use a specific search filter (Appendix 1). As this search strategy was not very specific, it yielded many results, which had to be screened in detail by two review authors. For the updated search, we integrated the search filter by Geersing 2012 to allow for a more specific strategy (Appendix 2).

We searched the following databases without applying any language restrictions in order to reduce language bias.

- MEDLINE via Ovid (searched 24 June 2019; Appendix 1; Appendix 2).
- Embase (searched 24 June 2019; Appendix 3).

We searched the following databases for ongoing trials.

- ClinicalTrials.gov (clinicaltrials.gov; searched 5 March 2020; Appendix 4).
- World Health Organization International Clinical Trials Registry Platform (WHO ICTRP) (searched 5 March 2020; Appendix 5).

Initially, we planned to include only models published after 1990. However, prognostic model studies are often conducted based on retrospective data, which in some cases included the analysis of blood samples frozen at diagnosis and analysed many years later. Thus, we decided not to limit our search strategy to date of publication. We did not screen conference proceedings, according to the section 'Types of prognostic models', where we stated that we would exclude conference abstracts based on limited information that would not allow us to assess risk of bias. In contrast to our planning (Skoetz 2016), we did not search the database of prognostic studies by the Prognostic Methods Group as it has not been developed. We did not search conference proceedings because of their limited information that would have complicated the inclusion and exclusion of studies as well as the risk of bias rating.

Searching other resources

As prespecified in the protocol (Skoetz 2016), we searched the following sources.

- Handsearching of references
 - References of all identified trials, relevant review articles and current treatment guidelines for further literature.
- Personal contacts
 - We contacted authors of relevant studies, study groups, experts and investigators from transplantation centres worldwide who are known to be active in the field for unpublished material or further information on ongoing studies.

In addition, and not reported in the protocol of this review, we tracked the citations of all included studies in October 2018 to identify possible additional validation studies of a developed model (Web of Science citation indexing).

Data collection and analysis

Selection of studies

During independent screening of titles and abstracts by two review authors (AA, LE, MT, NK, NS), we organised regular group discussions for congruency due to the novelty of the topic. The main discussion points can be found below. After title and abstract screening, full texts for all eligible studies were obtained and independently selected by two review authors (AA, LE, MT, NK, NS). Disagreements were solved by involving one or more additional review authors (Lefebvre 2019).

As recommended in the PRISMA statement (Moher 2009), we documented the total number of retrieved references and numbers of included and excluded studies in a flow chart.

Solving disagreements in study inclusion

The poor reporting and the fine line between the development of a prognostic model and prognostic factor (identification) studies made the screening of titles and abstracts and the final decision regarding the inclusion of studies challenging. During several group discussions, the following issues emerged repeatedly.

 As we changed our pre-planned limitation of the search to include studies from inception of the database, we obtained many studies labelled as "staging system." A staging system intends to describe the severity of a disease and, thus, indicate



some prognostic and therapeutic information. Two staging systems for CLL, Rai and Binet, are commonly used in clinical practice. However, the predicted survival of individuals with the same disease stage can be highly variable. We decided to exclude staging systems, as they are not used by clinicians any longer (except for Rai and Binet). Instead, we focused on new algorithms with a high chance to include more recently identified prognostic factors.

- We decided to exclude studies that build genetic signatures aiming to distinguish between good and poor prognosis of an individual, due to several reasons. First, the development of a genetic signature is often preceded by the identification of genetic markers on the same cohort from several hundred candidate markers, which introduces a high potential for overfitting of the algorithm to the development cohort (e.g. Houldsworth 2014). Second, genetic signatures can consist of a tremendous number of genes compared to the number of individuals included in the cohort (e.g. Ferreira 2014). Third, it is currently difficult to apply the algorithm of a genetic signature to an external cohort due to the complexity of the model.
- The fine line between a multivariable model, which includes several prognostic factors to prove their independence from each other, and a prognostic model was not always evident from the abstract. When an abstract mentioned a multivariable model and the formation of risk groups, we considered the full text of the paper. We then excluded papers that did not proceed to build a prognostic model or score explicitly.
- References were classified as an external validation study if
 the term "validation" was explicitly stated or the application
 of the prognostic model was clearly the main focus of the
 paper. We also included some publications where validation
 was not explicitly mentioned and performance measures were
 not reported, but where the authors put their focus on the
 application of one of the previously developed prognostic
 models (e.g. CLL-IPI V Reda 2017 (Milano cohort); CLL-IPI V Rigolin 2017 (Ferrera cohort)).

The decisions detailed above may seem subjective and were made in the case of this specific review. We would advise future authors who aim to evaluate prognostic models for a certain disease to clearly define the eligibility of different types of prognostic models at the protocol stage. This will help with objectivity throughout the screening process.

Data extraction and data management

Teams of two review authors (AA, LE, MT, NK, NS) independently extracted the data to enable assessment of applicability of the model for the review and the 'Risk of bias' assessment (see below). We contacted authors of individual studies for additional information, where required. We developed a standardised data extraction form containing the following items based on the CHARMS checklist (Moons 2014).

- · General information
 - Author, title, publication date, country, language, duplicate publications
- Source of data
 - E.g. cohort, case-control, randomised trial participants, or registry data
- · Participants

- Participant eligibility and recruitment method (e.g. consecutive participants, location, number of centres, setting, inclusion and exclusion criteria)
- Participant description
- o Details of treatments received
- Study dates
- · Outcomes to be predicted
 - o Definition and method for measurement of outcome
 - Was the same outcome definition (and method for measurement) used in all patients?
 - Was the outcome assessed without knowledge of the candidate predictors (i.e. blinded)?
 - Time of outcome occurrence or summary of duration of follow-up
- Candidate predictors
 - Number and type of predictors (e.g. demographics, patient history, physical examination, additional testing, disease characteristics, tumour markers)
 - Definition and method for measurement of candidate predictors
 - Timing of predictor measurement (e.g. at patient presentation, at diagnosis, at treatment initiation)
 - Were predictors assessed blinded for outcome, and for each other (if relevant)?
 - Handling of predictors in the modelling (e.g. continuous, linear, non-linear transformations or categorised)
- Sample size
 - Number of participants and number of outcomes/events
 - Number of outcomes/events in relation to the number of candidate predictors (events per variable)
- Missing data
 - Number of participants with any missing value (include predictors and outcomes)
 - o Number of participants with missing data for each predictor
 - Handling of missing data (e.g. complete-case analysis, imputation, or other methods)
- Model development
 - Modelling method (e.g. logistic, survival, neural networks, or machine learning techniques)
 - o Modelling assumptions satisfied
 - Method for selection of predictors for inclusion in multivariable modelling (e.g. all candidate predictors, preselection based on unadjusted association with the outcome)
 - Method for selection of predictors during multivariable modelling (e.g. full model approach, backward or forward selection) and criteria used (e.g. P value, Akaike Information Criterion)
 - Shrinkage of predictor weights or regression coefficients (e.g. no shrinkage, uniform shrinkage, penalised estimation)
- Model performance
 - Calibration (calibration plot, calibration slope, Hosmer-Lemeshow test) and discrimination (C-statistic, D-statistic, log-rank) measures with confidence intervals
 - Classification measures (e.g. sensitivity, specificity, predictive values, net reclassification improvement) and whether a priori cut-points were used
- Model evaluation



- Method used for testing model performance: development dataset only (random split of data, re-sampling methods, e.g. bootstrap or cross-validation, none) or separate external validation (e.g. temporal, geographical, different setting, different investigators)
- In case of poor validation, whether model was adjusted or updated (e.g. intercept re-calibrated, predictor effects adjusted, or new predictors added)
- Results
 - Final and other multivariable models (e.g. basic, extended, simplified) presented, including predictor weights or regression coefficients, intercept, baseline survival, model performance measures (with standard errors or confidence intervals)
 - Any alternative presentation of the final prediction models, e.g. sum score, nomogram, score chart, predictions for specific risk subgroups with performance
 - Comparison of the distribution of predictors (including missing data) for development and validation datasets
- · Interpretation and discussion
 - Interpretation of presented models (confirmatory, i.e. model useful for practice versus exploratory, i.e. more research needed)
 - Comparison with other studies, discussion of generalisability, strengths and limitations

References of model development studies, for which no external validation studies have yet been published, are summarised in Studies awaiting classification. These studies fit the formal inclusion criteria of this review. However, they have not yet been tested in any additional cohort, which renders summary of performance in external cohorts impossible. They cannot yet be evaluated regarding their clinical usefulness, and are therefore not yet described in detail.

Risk of bias and applicability assessment

Since the publication of the review protocol, a 'Risk of bias' tool specifically designed for prognostic model studies (Prediction model Risk Of Bias ASsessment Tool, PROBAST) was published (Moons 2019; Wolff 2019). Instead of the CHARMS checklist for critical appraisal of prognostic modelling studies as reported in our protocol, we used this new tool recommended by the Cochrane Prognosis Methods Group to assess the risk of bias of the individual prediction models investigated in the included primary studies. Teams of two review authors (AA, LE, MT, NK, NS) independently assessed the risk of bias and applicability for each study. We assessed only validation studies with the outcome that a model was developed for, in line with meta-analyses, hence PROBAST assessments reflect only the outcome of interest and not any additional outcomes reported in the same study.

After classifying each study into one of the three categories (model development with or without external validation in the same publication and external validation study of a previously developed model only), we assessed risk of bias according to the following four PROBAST domains (once per developed or validated model and per outcome).

- Participants
- Predictors
- Outcome

Analysis

We answered signalling questions within each domain with one out of five options ('yes', 'probably yes', 'probably no', 'no', 'no information'); 'yes' always implied the absence of a potential bias-generating aspect. We rated domain-level 'Risk of bias' assessments using one of the following three options.

- Low risk of bias: if the criterion is adequately fulfilled in the study, i.e. the study is at a low risk of bias for the given domain.
- High risk of bias: if the criterion is not fulfilled in the study, i.e. the study is at high risk of bias for the given domain.
- Unclear risk of bias: if the study report does not provide sufficient information to allow for a clear judgement or if the risk of bias is unknown for one of the domains listed above.

PROBAST additionally requires the judgement of the applicability of the model to the research question. The assessment occurs per domain (for the first three domains only) with the following response options: 'low concern regarding applicability', 'high concern regarding applicability' and 'unclear concern regarding applicability' (equivalent to the categories for risk of bias).

Based on these domain-level judgements, we established an overall judgement per study based on PROBAST guidance for risk of bias:

- Low risk of bias: if a prediction model evaluation is judged as low on all domains relating to bias or applicability.
- High risk of bias: if an evaluation is judged as high for at least one domain.
- Unclear risk of bias: if the prediction model evaluation is unclear in one or more domains and was rated as low in the remaining domains

When information for a complete judgement was missing, we contacted the corresponding authors via email to request additional information to be able to make sound judgements on the 'Risk of bias' assessment. According to PROBAST, models that were developed, but not (yet) externally validated can be classified as 'high risk of bias', with the exception of extremely large samples (Wolff 2019). Due to the novelty of PROBAST and the subjectivity of ratings, we devoted a small section of the results to present our experience and agreements. Two authors of PROBAST (RW, KGMM) are also authors in our review and RW was involved in group discussions addressing disagreements between ratings.

Solving disagreements regarding PROBAST ratings

Due to the novelty of the 'Risk of bias' tool (PROBAST), we arranged frequent team meetings to discuss disagreements and discrepancies regarding ratings between the six authors who extracted data and assessed risk of bias and applicability. For reasons of transparency, we report the following challenges and our agreements.

 The relevant observation time varies between diseases. For an indolent neoplasm like CLL, the clinicians in our team considered a median follow-up of five years as appropriate for OS in a cohort with a normal case mix (i.e. no limitation to patients with late stage or high-risk disease such as patients with del(17p) only).



- The signalling question for the participant domain asks whether all inclusions and exclusions of participants were appropriate (item 1.2). Some publications listed missing values as part of the exclusion criteria. Missing values are also treated in the analysis domain (items 4.3 and 4.4). To avoid duplicate rating of the same issue, we decided to rate the risk of bias as high for the participant domain when complete data were required for eligibility, and rated the risk of bias as high for the analysis domain when the number and characteristics of the background sample were described but persons were excluded from analysis due to missing values.
- Some publications did not report eligibility criteria and/or recruitment period. We rated these studies as unclear not only for risk of bias, but also for concerns of applicability, because we could not be sure if the included individuals matched our review question and a current application of the model.
- When a publication lacked information necessary to answer a signalling question, but other available information could be used as an indication, we made assumptions regarding the answer (probably yes, probably no) to avoid the option 'no information'.
- For the CLL-IPI, we frequently encountered the problem that one
 predictor was replaced by a proxy predictor in the analysis of
 an external validation study. We assumed that this may happen
 in clinical practice, and included the external validation study
 to validate the CLL-IPI, although strictly speaking, they did not
 apply the original model. We rated the concern for applicability
 as unclear in the domain predictor.
- Some models developed to predict TFS include predictors also used for treatment indication, which means that the predictors are not excluded from the outcome definition (PROBAST item 3.3; e.g. O-CLL1 model: Rai staging is included in the prognostic score, and is, in combination with other characteristics, a factor for treatment indication). This may lead to an overestimation of the association between predictor and outcome, which is called incorporation bias (Moons 2019). We answered the corresponding PROBAST item 3.3 with 'no' and considered the risk of bias to be high for the outcome domain. We decided that all corresponding external validations, which validate a model that include a predictor also used for treatment indication, are also considered to contain a high risk of bias.
- The PROBAST guidance for item 4.1, which states that a study should include at least 20 events per candidate predictor for development studies and 100 events for validation studies is recommended for regression analysis with a binary outcome. The models included in our systematic review are mainly based on Cox proportional hazard models (Moons 2019). We therefore considered a cohort with more than 100 events to be an appropriate sample size for development and external validation samples (Collins 2016).
- The testing of model assumptions (PROBAST item 4.6) was very rarely reported. We did not let this item influence our judgement regarding the analysis domain because, by default, all except one study would have been considered to be of unclear risk of bias. We assumed that authors tested the pre-conditions before using their analysis type. However, we would advise authors of primary studies to improve reporting of the entire model development process.

Measures of prediction model performance

In contrast to 'classical' meta-analysis focusing on treatment efficacy as the parameter of interest, there is no single recommended methodology to meta-analyse the predictive performance of prognostic models (Debray 2012; Debray 2014). Hence, we extracted the reported predictive performance measures for each studied model from the development and validation studies, notably the model's discrimination and calibration in the development cohort (i.e. apparent performance, the performance measures after internal validation) and the performance found in a model's external validation.

Calibration refers to the accuracy of the predicted risk probabilities, which means the agreement between estimated and observed number of events in a cohort. A model might predict more events than the number of observed events, i.e. there is overestimation of the number of events. In contrast, a model can predict fewer events than observed meaning that the number of events is underestimated. In both situations, the model is miscalibrated. The use of a miscalibrated model in clinical practice can misguide clinical decisions based on such models (Van Calster 2019). Calibration can be presented as a calibration plot (expected probabilities plotted against observed outcome frequencies), as ratios between observed and expected number of events or outcome frequencies (O:E ratios) or calibration table (Debray 2017).

Discrimination refers to the ability of a prediction model to differentiate between those who do or do not experience the outcome event. A model has perfect discrimination if the predicted risks for all individuals who develop the outcome are higher than those for all individuals who do not experience the outcome. Discrimination is commonly estimated by the socalled concordance index (c-statistic, also sometimes called cindex). The c-statistic reflects the probability that for any randomly selected pair of individuals, one with and one without the outcome, the model assigns a higher probability to the individual with the outcome. The c-index can range from 0 to 1, with 1 indicating perfect discriminative ability and 0.5 meaning the model's predictions equal chance. The c-index is identical to the area under the receiver-operating characteristic (ROC) curve for models with binary endpoints, and can be generalised for time-toevent (survival) models accounting for censoring (Debray 2017).

Both measures are needed to assess the performance of a prognostic model.

Unit of analysis issues

Ideally, a prognostic model is developed and tested on specially designed, prospectively followed cohort studies consisting of a representative case mix. However, more often we identified models or validations derived from retrospective data or data from randomised controlled trials (RCTs). Data from these populations are used for several purposes and models. Where a model was developed and externally validated in the same publication, we compared the institution and year of inclusion of individuals to assure independence of the cohorts. We also compared the institution and year of inclusion of all external validation cohorts belonging to one developed model to avoid overlaps in participants. When the same cohort was used to validate two or more different prognostic models, we considered them as separate studies. When RCT data were used, the randomisation of



individuals was not considered by the authors of prognostic model studies.

Dealing with missing data

When data were missing, we requested additional information from the original investigators. When confidence intervals for measures of discrimination were still missing after we contacted the author, we calculated these according to a method proposed by Newcombe (Debray 2017; Debray 2018a; Newcombe 2006).

Investigation/description of heterogeneity

For the c-statistic, we used the between-study standard deviation $(tau - \tau)$ to quantify possible heterogeneity. As we did not summarise O:E ratios to pool calibration, we merely visually inspected differences in survival per subgroup between cohorts. We planned to investigate and discuss clinical and statistical heterogeneity and design aspects of included studies mentioned in the section 'Data extraction and data management' based on subgroup analysis. We were unable to perform prespecified subgroup analysis based on diagnostic criteria due to the low number of studies with clear distinction between criteria.

Discussing reporting deficiencies

It is widely recommended that all prognostic models assess and report calibration and discrimination (Collins 2015; Moons 2015). However, it is known from numerous systematic reviews on methodological conduct and reporting of prognostic models in various disciplines that calibration is rarely reported and when it is reported (Heus 2018), it is done quite poorly. Hence, we also evaluated reporting deficiencies.

Methods and reporting in prognostic research often do not follow current methodological recommendations, limiting retrieval, reliability and applicability of these publications (Bouwmeester 2012; Peat 2014). There are some indications that prognosis research is cluttered with false-positive studies which would not have been published if the results were negative. Moreover, studies evaluating development studies of prognostic models are not prospectively registered, and usually no protocol is published (Peat 2014). Therefore, it is difficult to assess publication bias. We used sensitive search strategies to increase retrieval (Geersing 2012).

Data synthesis

Data synthesis and meta-analysis approaches

In the protocol, we explained that we would pool the performance measures for calibration and discrimination. During the review process however, new methodological developments became available and we chose to adopt them, as follows. Although we assessed both model development and external validations for risk of bias, we performed meta-analysis only for validation studies, as there was only one development study per model.

- When a particular model has been validated at least three times for the primary outcome, we applied a random-effects model for pooling the logit transformation of the discrimination measure (the c-statistic) using the meta-analysis packages 'metamisc' and 'metafor' in the R statistical language (Debray 2014; Debray 2018b; Viechtbauer 2010).
- We pooled only validation studies with the outcome that a model was developed for (e.g. we did not meta-analyse

- validation studies that tested a model to predict TFS if the model was developed for OS).
- We used random-effects meta-analysis, because validation studies typically differ regarding several parameters, including patient characteristics and design.
- Where the c-statistic was not reported, we did not estimate
 it. When the c-statistic was provided without measures of
 uncertainties, we calculated the standard error and confidence
 intervals based on the P value or the combination of sample size
 and number of events, if available, according to Newcombe and
 colleagues (Debray 2017; Debray 2018a; Newcombe 2006).

We planned to summarise the measures of calibration. Unfortunately, calibration measures and the expected survival, both in development and external validation cohorts, were rarely reported. Instead, many studies reported the observed outcome frequency per subgroup at a specific time point. Though it is possible to estimate approximate O:E ratios from the reported observed survival per risk category and probabilities reported in the model development study (for scores), in collaboration with the Cochrane Prognosis Methods Group, we decided against this estimation because it is merely an approximate indication for the calibration of a model. For time-to-event outcomes, we would need to account for censoring (Debray 2018a), which was not possible based on the limited reporting of the studies. Instead, we illustrated the survival per risk category for each model with sufficient data graphically. We pooled survival per risk category for all external validations of one model and reported a prediction interval. The 95% prediction interval (95% PI) is an estimate of the range in which a future average survival frequency in a new validation study of the prognostic model will fall with a 95% probability.

Subgroup analysis

We planned to investigate whether the change of definition of CLL over time has affected the performance of a model (i.e. Cheson 1996; Hallek 2008). We did not perform this analysis because the number of studies that reported a clear distinction between the definitions was too low.

Sensitivity analysis

During the review process, and before data extraction and analysis, we decided to conduct sensitivity analysis based on the 'Risk of bias' rating, if a sufficient number of validation studies were available per model.

We decided post hoc to conduct the following model-specific meta-analyses.

- Test the effect of studies that reported the area under the curve (AUC) instead of the c-statistic (which we meta-analysed together). These two measures for discrimination are the same for binary outcomes, but may differ for time-to-event data depending on the time point chosen to calculate the AUC.
- Explore the effect of the estimation of the 95% CI for the c-statistic. Estimating confidence intervals can introduce imprecision, therefore we aimed to explore the extend of the changes between reported and estimated 95% CIs.
- For the CLL-IPI, several external validation studies used a proxy prognostic factor for TP53 deletion or mutation, the cytogenetic aberration based on FISH, del(17p). Although we expected the effect to be small as the concordance between the two



predictors is larger than 90% (Dicker 2009), using a proxy can reduce predictive performance.

Rating the certainty of evidence and summary of findings

Originally, we planned to use the most up-to-date GRADE guidance for this systematic review. However, no GRADE guidance for grading the certainty of results from meta-analysis of prognostic models yet exists. Hence, for this review, we decided to refrain from applying

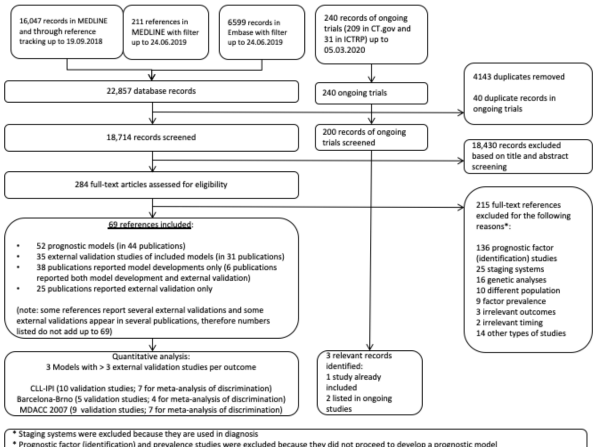
RESULTS

Description of studies

Results of the search

Our literature search (24 June 2019; MEDLINE and Embase) resulted in 22,857 potentially relevant references related to prognostic models for newly-diagnosed untreated individuals with chronic lymphocytic leukaemia (CLL). Of these, we removed 4144 duplicates, leaving 18,713 references for title and abstract screening. We identified 284 full-text references, which might fulfil our predefined inclusion criteria, including two references identified by reference and citation tracking. Of these, we excluded 214 references. We documented the overall number of screened, included and excluded references in a PRISMA flow diagram (Figure

Figure 1. Study flow diagram



^{*} Prognostic factor (identification) and prevalence studies were excluded because they did not proceed to develop a prognostic model

Other types of excluded studies: risk factor identification, diagnostic accuracy studies, reviews, commentary on excluded study

This led to a total of 70 included references. Of these, 38 reported model developments only, six publications reported both model development and external validation(s), and 25 publications reported external validation only. In total, we identified 52 prognostic models (in 44 publications), and 35 external validation studies of included models that predicted the primary outcome of one of the included developed models (in 31 publications). Several of these publications offered secondary analyses with another

outcome. These are only listed in this review if the primary outcome was not reported.

Out of the 52 developed models, 12 models were externally validated at least once either in the primary publication or as described in an additional publication (for an overview, see Appendix 6). Forty-one models did not undergo any external validation.



The search of databases for ongoing trials resulted in 240 records (ClinicalTrials.gov: 209 and ICTRP: 31; 5 March 2020). Of these, 40 records were duplicates. We identified three records that referred to the potential use of developing a prognostic model. One cohort had already been included from the database search (O-CLL-GISL, development of a model and several validations, e.g. O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL)). One study is currently recruiting (NCT03436524), and one mentioned the aim to develop a staging system, however, it does not refer to any relevant publication yet (NCT00275054).

It is important to note the difference between 'reference' and 'study': one reference of a model development study may in addition to the model development also report one or several external validation studies of their model in independent cohorts. Furthermore, some references of external validations may report and compare the performances of several prognostic models in their cohort. Several publications used the same dataset to validate the same model(s).

References of model development studies without any external validation studies are summarised in Studies awaiting classification and Appendix 7.

Included studies

We described the characteristics of included studies per model and per predicted outcome. The numbers of external validations belonging to each developed model and outcome may not correspond to the number of included references. We defined a study as a validation of one model in one independent cohort, which implies that the validation of two different models in the same cohort are counted as two separate studies for the description in this review. In some publications, one model was externally validated in several cohorts, while other publications validated several models in one cohort. We presented the number of participants included in analysis instead of the number of participants in the study sample or complete registry as, often, missing values were reported to be an exclusion reason. We only included external validation studies that predicted the same outcome that the prognostic model was developed for.

Models with meta-analysis (> 3 external validations)

- CLL-IPI (OS, CLL-IPI D Bahlo 2016 (development cohort))
- Barcelona-Brno model (OS, Barcelona-Brno D Delgado 2017 (Barcelona cohort))
- MDACC 2007 (OS, MDACC 2007 D Wierda 2007 (MDACC))

Models without meta-analysis (1-3 external validations)

- GCLLSG model (OS, GCLLSG D Pflug 2014 (GCLLSG))
- Rossi model (OS, Rossi D Rossi 2013 (Italian cohort))

- Stephens (OS, Stephens OS D Stephens 2015 (Ohio cohort))
- Baliakas model (TFS, Baliakas D Baliakas 2019 (multicentre))
- GIMEMA model (PFS, GIMEMA D Molica 2005 (GIMEMA cohort))
- MDACC 2011 (TFS, MDACC 2007 D Wierda 2007 (MDACC))
- Morabito model (TFS, Morabito D Morabito 2009 (Italian cohort))
- O-CLL1 model (TFS, O-CLL-1 D Gentile 2016 (O-CLL-1-GISL))
- Stephens model (TFS, Stephens TFS D Stephens 2015 (Ohio cohort))

Models with more than three external validation studies per outcome

Prognostic models for the prediction of OS (with meta-analysis, > 3 external validations)

CLL-IPI (CLL-IPI D - Bahlo 2016 (development cohort))

The CLL-IPI model predicts OS and was derived from combined data from eight phase three RCTs conducted in multiple countries (France, Germany, Poland, UK, USA), with a total of 1799 individuals included for the model development. The recruitment period for the trials lasted from 1997 to 2009 (CLL-IPI D - Bahlo 2016 (development cohort)). The median follow-up time was 79.9 months (interquartile range (IQR) 79.9 to 101.4 months). All disease stages were included; most individuals were classified as Rai I/ II or Binet B stage (50% and 41% respectively). The final model includes five predictors: TP53 status, IgHV mutational status, serum B2-microglobulin, clinical stage (Rai or Binet), and age. The original weighting of predictors was simplified to present as a risk score for easier use in clinical practice. We identified 11 external validation cohorts (CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014); CLL-IPI V -Bahlo 2016 (SCAN cohort); CLL-IPI V - Da Cunha-Bang 2016 (Danish cohort); CLL-IPI V - Delgado 2017 (Barcelona cohort); CLL-IPI V -Gentile 2016 (Italian cohort); CLL-IPI V - Molica 2016 (O-CLL1-GISL); CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort); CLL-IPI V - Rani 2018 (Indian cohort); CLL-IPI V - Reda 2017 (Milano cohort); CLL-IPI V - Rigolin 2017 (Ferrera cohort); CLL-IPI V - Zhu 2018 (Chinese cohort)), with 10 validations for OS and eight for TFS. The validation studies for OS included a total of 5485 individuals with available data presenting between 1983 and 2016. Two relevant validation studies lacked information regarding the recruitment period (CLL-IPIV - Delgado 2017 (Barcelona cohort); CLL-IPIV - Rani 2018 (Indian cohort)). Due to lack of reporting, only 3307 individuals with 917 deaths from seven external validation studies were included in the meta-analysis of the discrimination (CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014); CLL-IPI V - Bahlo 2016 (SCAN cohort); CLL-IPI V -Delgado 2017 (Barcelona cohort); CLL-IPI V - Gentile 2016 (Italian cohort); CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort); CLL-IPI V -Rani 2018 (Indian cohort); CLL-IPIV - Zhu 2018 (Chinese cohort)). We provide a summary of the main characteristics of studies regarding the CLL-IPI in Figure 2.



Figure 2. CLL international prognostic index - summary of characteristics of included studies

							Characteristi	ics of included stud	ies		
Mo	odel	Study	recruitment	country	number of events / number persons	age	gender	stage	treatment	design	follow-up
L-IPI model verall survival)	Development	Bahlo 2016 (Development cohort)	1997 - 2009	France, Germany, Poland, UK, USA, etc.	462 events 1214 persons (train) 243 events 585 persons (test)	median 61 years (range: 27 - 86 years)	70% male	Rai 0: 14% Rai I/II: 50% Rai III/IV: 36%	690 F 1017 FC 113 CC 475 FCR 81 FCCAM 479 CHR	combined RCT data	median 79.9 months, IQ 79.9 - 101.4
		Bahlo 2016 (Mayo clinic 2001-2014)	2001 - 2014	USA	144 events 838 persons	≤ 65 years: 533 persons > 65 years: 305 persons	67.8% male	Rai 0: 57.4% Rai I-II: 38.5% Rai III-IV: 4.1%	168 purine analogs w/o MoAbs 37 Alkylator-based Chemoimmunotherapy 11 Alkylator w/o MoAbs 97 antibody only 7 RTK inhibitors 6 other	prospective cohort	median 63.2 months, IQ 30.2 - 91.8
		Bahlo 2016 (SCAN cohort)	1999 - 2002	Denmark, Sweden	215 events 416 persons	≤ 65 years: 242 persons > 65 years: 174 persons	62.7% male	Binet A: 78.1% Binet B: 16.6% Binet C: 5.3%	NR	unclear	median 151 months, IQF 124.8 - 163.0
		Delgado 2017 (Barcelona cohort)	NR	Spain	212 events 524 persons	median 62 years (range: 22 - 93 years)	60% male	Rai 0: 62%; Rai I- IV: 38%	83 FCR or similar 82 purine alanogs w/o MoAbs 86 received alkylating agents 23 received others	retrospective cohort	median 99.6 months (range 1 - 456 months)
	Meta-analysed c-statistic	Gentile 2016 (Italian cohort)	1985 - 2015	Italy	174 events 858 persons	median 65.5 years (range: not reported)	56.2% male	Rai 0: 58.4% Rai I: 19.8% Rai II: 15.2% Rai III: 1.5% Rai IV: 5.1%	130 received chemotherapy 174 received chemo- immunotherapy	retrospective cohort	median 69.6 months, range not reported
		Muñoz-Novaz 2018 (Spanish cohort)	1989 - 2013	Spain	47 events 258 persons	median 65.7 years (IQR: 55.2 - 73.5)	62.8% male	Rai 0: 58.9%; Rai I-IV: 41.1%	NR	retrospective cohort	median 68 months (rang 3 - 277 months)
		Rani 2018 (Indian cohort)	NR	India	86 events 198 persons	median 60 years (range: not reported)	77% male	Rai 0: 14.6%; Rai I: 21.2%; Rai III: 34.9%; Rai III: 14.1%; Rai IV: 15.2%	62 CHL 56 rituximab-based therapy 20 other therapies	NR	median 40.5 months (range 1 - 215 months)
		Zhu 2018 (Chinese cohort)	2002 - 2017	China	46 events 215 persons	median 60 years (range 16 - 85 years)	66.5% male	Rai 0: 16.7% Rai I-II: 54.0% Rai III-IV: 29.3%	147 treated with benzodiazepine FCR bendamoxetine R ReFFP+HDMP	retrospective cohort	median 48 months, (range: 1 - 192 months)
		Da Cunha-Bang 2016 (Danish cohort)	2008 - 2015	Denmark	249 events 1514 persons	median 69 years (range: not reported)	60% male	Binet A: 80% Binet B/C: 20%	NR	prospective cohort	median 38.4 months, range not reported
		Molica 2016 (O-CLL1- GISL)	2007 - NR	Italy	91 events (TFS) 337 persons	median 61 years (range: 33 - 70 years)	57.2% male	Rai 0: 77.8% Rai ≥ I: 22.2%	NR	prospective cohort	median 42 months, 2038 person-years
1	Narrative	Reda 2017 (Milan cohort)	1983 - 2016	Italy	NR events 329 persons	median 65 years (range: 32 - 91 years)	57% male	Rai 0-1: 74%; Rai II-IV: NR	190 CHL 108 FCR 12 O-Benda 44 BR 65 alemtuzumab 29 ibrutinib or i delalisib	retrospective cohort	median 144 months (range 0 - 360 months)
		Rigolin 2017 (Ferrera cohort)	2006 - 2016	Italy	NR events 335 persons	median 68.7 years (range: 33 - 96 years)	57.9% male	Binet A: 77.9% Binet B: 14.3% Binet C: 7.8%	114 chemotherapy	retrospective cohort	NR

Abbreviations: NR: not reported; CHL: chiorambuci; F: fluradabine; CC: cyclophosphamide + cladribine; FC: fluradabine + cyclophosphamide; FCR: fluradabine + cyclophosphamide + ritusimab; FCAM: FC+ alemtuzumab; MoAbs: monoclonal antibody; ritusimab; Fresh frozen olsama; EHDMP; high-dose methylopedisolone

Barcelona-Brno model (Barcelona-Brno D - Delgado 2017 (Barcelona cohort))

The Barcelona-Brno model predicts OS and was derived from data from a retrospective, single-centre cohort study conducted in Spain (Barcelona-Brno D - Delgado 2017 (Barcelona cohort)). TFS was reported as a secondary outcome. Model development included 524 individuals. The recruitment period was not reported. The median follow-up time was 99.6 months, ranging from 1 to 456 months. All disease stages were included; most individuals were Rai 0 or Binet A stage (62% and 83% respectively). The model aimed to simplify the CLL-IPI described below, therefore only combinations of the factors included in the CLL-IPI were tested. The final model included three predictors: IgHV mutational status, and the genomic aberrations del(17p), and del(11q). We identified six external validation cohorts (Barcelona-Brno V - Delgado 2017 (Brno cohort); Barcelona-Brno V - Gentile 2017 (Italian & Mayo);

Barcelona-Brno V - Molica 2017 (O-CLL1-GISL); Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.); Barcelona-Brno V - Rani 2018 (Indian cohort); Barcelona-Brno V - Reda 2017 (Milan cohort)), with five validations for OS and six for TFS. The validation studies for OS included a total of 2501 individuals presenting between 1983 and 2016. Two validation studies lacked information regarding the recruitment period (Barcelona-Brno V - Delgado 2017 (Brno cohort); Barcelona-Brno V - Rani 2018 (Indian cohort)). Due to lack of reporting, only 1755 individuals with 416 deaths from three studies were included in meta-analysis of discrimination (Barcelona-Brno V - Delgado 2017 (Brno cohort); Barcelona-Brno V - Gentile 2017 (Italian & Mayo); Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.); Barcelona-Brno V - Rani 2018 (Indian cohort)). We provide a summary of the main characteristics of studies regarding the Barcelona-Brno model in Figure 3.



Figure 3. Barcelona-Brno score - summary of characteristics of included studies

			l				Characteristi	cs of included stud	ies		
Mo	del	Study	recruitment	country	number of events / number persons	age	gender	stage	treatment	design	follow-up
Barcelona-Brno model (overall survival)	Development	Delgado 2017 (Barcelona cohort)	NR	Spain	212 events 524 persons	median 62 years (range: 22 - 93 years)	60% male	Rai 0: 62%; Rai I- IV: 38%	83 FCR or similar 82 purine alanogs w/o MoAbs 86 received alkylating agents 23 received others	retrospective cohort	median 99.6 months (range 1 - 456 months)
		Delgado 2017 (Brno cohort, CZ)	NR	Czech Republic	158 events 417 persons	median 62 years (range: 32 - 85 years)	66% male	Rai 0: 41%; Rai I- II: 43%; Rai III- IV: 16%	125 FCR or similar 41 purine analogs w/o MoAbs 53 alkylating agents 57 received others	NR	median 59 months (range 3 - 330 months)
Meta-analysec c-statistic	Meta-analysed c-statistic	Gentile 2017 (Italian multicentre and Mayo cohort)	1985 - 2015	Italy & USA	283 events 1299 persons	median 63 years (range: 27 - 92 years)	61.3% male	Rai 0: 57.9%, Rai I: 28.6% Rai II: 8.6%, Rai III: 1.5%; Rai IV: 3.5%	194 chemotherapy 316 chemoimmunotherapy	retrospective cohort	median 82 months (range 3 - 330 months)
		Muñoz-Novaz 2018 (Spanish cohort)	1989 - 2013	Spain	47 events 258 persons	median 65.7 years (IQR: 55.2 - 73.5)	62.8% male	Rai 0: 58.9%; Rai I-IV: 41.1%	137 received therapy, unclear which therapy	retrospective cohort	median 68 months (range 3 - 277 months)
		Rani 2018 (Indian cohort)	NR	India	86 events 198 persons	median 60 years (range: not reported)	77% male	Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%	62 CHL 56 rituximab-based therapy 20 other therapies	NR	median 40.5 months (range 1 - 215 months)
		Molica 2017 (O-CLL1- GISL)	2007 - NR	Italy	91 events 337 persons	median 61 years (range: 33 - 70 years)	57.2% male	Rai 0: 77.8%; Rai I-II: 22.2%	NR	prospective cohort	median 42 months (range 1 - 82 months)
	Narrative	Reda 2017 (Milan cohort)	1983 - 2016	Italy	NR events 329 persons	median 65 years (range: 32 - 91 years)	57% male	Rai 0-1: 74%; Rai II-IV: NR	190 CHL 108 FCR 12 O-Benda 44 BR 65 alemtuzumab 29 Ibrutinib or I delalisib	retrospective cohort	median 144 months (range 0 - 360 months)

MDACC 2007 (Wierda 2007) (MDACC 2007 D - Wierda 2007 (MDACC))

The MDACC 2007 model predicts OS and was derived from data from a retrospective, single-centre cohort study conducted in the USA (MDACC 2007 D - Wierda 2007 (MDACC)). A total of 1674 individuals were included for the model development. The recruitment period for the study lasted from 1981 to 2004. The median follow-up time was 58.8 months (95% confidence interval (CI) 55.2 to 61.2 months). All disease stages were included; most individuals were classified as Rai I or Binet A stage (739/1674 and 1019/1674 respectively). The final model included six predictors: age, absolute lymphocyte count, serum B2-microglobulin, nodal groups, Rai stage, and gender. The publication reports both the formula to be used in combination with a nomogram, and a simplified index score. We identified 10 external cohorts (MDACC 2007 V - Trajkova 2013 (Macedonia); MDACC 2007 V - Bulian 2011 (Italian-Swiss); MDACC 2007 V - Gentile 2014 (Italian cohort); MDACC 2007 V - Gentile 2016 (Mayo cohort); MDACC 2007 V - González Rodríguez (Cabueñes); MDACC 2007 V - Molica 2010 (GIMEMA cohort); MDACC 2007 V -Molica 2015 (O-CLL1-GISL); MDACC 2007 V - Muñoz-Novas 2018

(Spanish cohort); MDACC 2007 V - Pflug 2014 (3 RCTs); MDACC 2007 V - Rani 2018 (Indian cohort)), with eight validations for OS and six for TFS. Seven studies used the index score and three additionally calculated the formula to predict OS. For one study, it was unclear if the index score or the formula were used. The eight cohorts for which OS was predicted included 6928 individuals presenting between 1983 and 2013. One validation study lacked information regarding the recruitment period (MDACC 2007 V - Rani 2018 (Indian cohort)). Only the simplified index score was sufficiently externally validated to be summarised in meta-analysis. In total, 5127 individuals with 994 deaths from seven studies remained for inclusion in the meta-analysis of discrimination (MDACC 2007 V -Bulian 2011 (Italian-Swiss); MDACC 2007 V - Gentile 2014 (Italian cohort); MDACC 2007 V - Gentile 2016 (Mayo cohort); MDACC 2007 V - González Rodríguez (Cabueñes); MDACC 2007 V - Muñoz-Novas 2018 (Spanish cohort); MDACC 2007 V - Pflug 2014 (3 RCTs); MDACC 2007 V - Rani 2018 (Indian cohort)). We provide a summary of the main characteristics of studies regarding the MDACC 2007 model in Figure 4.



Figure 4. MDACC 2007 index score - summary of characteristics of included studies

							Characteristi	cs of included stud	lies		
М	odel	Study	recruitment	country	number of events / number persons	age	gender	stage	treatment	design	follow-up
ACC 2007 del erall survival)	Development	Wierda 2007 (Development cohort)	1981 - 2004	USA	437 events 1617 persons	median 58 years (range: 0 - 90 years)	1029 male	Rai 0: 469 Rai I: 739 Rai II: 235 Rai III: 93 Rai IV: 127	NR	retrospective cohort	median 58.8 months, 95% CI: 55.2 - 61.2
		Bulian (Italian-Swiss cohort)	1996 - 2011	Italy, Switzerland	151 events 1037 persons	median 65 years (range: 21 - 94 years)	58.2% male	Binet A: 74.6% Binet B: 18.4% Binet C: 6.9%	306 chemotherapy 129 chemoimmunotherapy 42 FCR 49 FR 38 R-other 40 with missing data	retrospective cohort	median 63.6 months, range not reported
		Gentile 2014 (Italian multicentre cohort)	1983 - 2013	Italy	277 events 1502 persons	median 67 years (range: not reported)	55.7% male	Rai I: 21.4% Rai I: 21.4% Rai II: 15% Rai III: 2.6% Rai IV: 4.5%	337 chemotherapy 142 chemo-immunotherapy	retrospective cohort	median 68.4 months, range not reported
		Gentile 2016 (Mayo cohort 2001-2008)	2001 - 2008	USA	114 events 506 persons	median 62.5 years (range: 36 - 89 years)	68.6% male	Rai 0: 62.5% Rai I: 28.9% Rai II: 5.1% Rai III: 1.2% Rai IV: 2.4%	NR.	prospective cohort	median 86.4 months range not reported
	Meta-analysed	González Rodríguez 2009 (Cabueñes hospital cohort)	1997 - 2007	Spain	76 events 257 persons	median 71.7 years (range: 42 - 94 years)	NR	Binet A: 76.8% Binet B: 14.8% Binet C: 8.4%	NR	retrospective cohort	NR
	index score	Muñoz-Novaz 2018 (Spain multicentre cohort)	1989 - 2013	Spain	92 events 483 persons	median 67 years (range: 25 - 90 years)	64.2 % male	Rai 0-II: 93.8% Rai III-IV: 6.2%	NR	retrospective cohort	median 46 months, ra not reported
		Pflug 2014 (German CLL study group cohort)	1997 - 2006	Australia, Austria, Belgium, Czech Republic, Denmark, France, Germany, Israel, Italy, New Zealand, Spain	180 events 1144 persons	median 60 years (range: 30 - 81 years)	68.4% male	Rai 0: 20.8% Rai I: 18.4% Rai II: 37.9% Rai III: 8.6% Rai IV: 14.3%	early F FC FCR	combined RCT data	median 63.4 months range not reported
		Rani 2018 (Indian cohort)	NR	India	86 events 198 persons	median 60 years (range: not reported)	77% male	Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%	62 CHL 56 rituximab-based therapy 20 other therapies	NR	median 40.5 months (range 1 - 215 month
		Molica 2010 (GIMEMA cohort 1991-2000)	1991 - 2000	Italy	NR events 310 persons	median 64 years (range: not reported)	56.1% male	Rai 0: 67% Rai I-III: 32.9% Rai III/IV: 0%	NR	retrospective cohort	median 41.5 months range not reported
	Narrative	Molica 2015 (O-CLL1- GISL)	2007 - NR	Italy	91 events (TFS) 337 persons	median 61 years (range: 33 - 70 years)	57.1% male	Rai 0: 77.5% Rai I-II: 22.5%	NR	prospective cohort	median 42 months, r not reported
I	merrauve	Trajkova 2013 (Macedonian cohort)	2011 - 2013	Macedonia		median 64.8 years (range: 47 - 78 years)	63% male	Rai 0: 54% Rai I: 5% Rai II: 32% Rai III: 5% Rai IV: 4%	NR	unclear	NR

Prognostic models for the prediction of PFS or TFS (with meta-

analysis, > 3 external validations)

We did not identify any prognostic model development studies that were externally validated more than three times for the outcomes PFS or TFS.

Models with one to three external validation studies per outcome

Prognostic models for the prediction of OS (without meta-analysis, 1 to 3 external validations)

GCLLSG model (Pflug 2014) (GCLLSG D - Pflug 2014 (GCLLSG))

The GCLLSG model predicts OS and was derived from combined data from three RCTs conducted in multiple countries (Australia, Austria, Belgium, Czech Republic, Denmark, France, Germany, Israel, Italy, New Zealand and Spain), with a total of 2007 individuals

included for the model development (GCLLSG D - Pflug 2014 (GCLLSG)). The recruitment period for the trials lasted from 1997 to 2006. The median follow-up time was 63.4 months; ranges were not reported. All disease stages were included; most individuals were Rai II or Binet A stage (37.9% and 42.6%, respectively). The final model included eight predictors: gender, age, Eastern Co-operative Oncology Group Performance Status (ECOG PS), del17p, del11q, IgHV mutational status, serum thymidine kinase, and serum B2-microglobulin. We identified three external validation cohorts, with one validation for OS (676 individuals, GCLLSG V - Pflug 2014 (Mayo cohort)), and two for TFS (GCLLSG V - Molica 2015 (O-CLL1-GISL); GCLLSG V - Rani 2018 (Indian cohort)). All validation studies lacked information regarding the recruitment period. We provide a summary of the main characteristics of studies regarding the GCLLSG model in Figure 5.



Figure 5. GCLLSG model - summary of characteristics of included studies

Characteristics of	fincluded studies	s for the GCLLSG ext	ernally validate	d model								
							Characteristics of included studies					
Мо	del	Study	recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up	
GCLLSG model (overall survival)	Development	Pflug 2014 (Development cohort)	1997 - 2006	Australia, Austria, Belgium, Czech Republic, Denmark, France, Germany, Israel, Italy, New Zealand, Spain	180 events 1144 persons	median 60 years (range: 30 – 81 years)	68.4% male	Rai 0: 20.8% Rai I: 18.4% Rai II: 37.9% Rai III: 8.6% Rai IV: 14.3%	early F FC FCR	combined RCT data	median 63.4 months, range not reported	
		Molica 2015 (O-CLL1- GISL)	2007 - NR	Italy	91 events 337 persons	median 61 years (range: 33 - 70 years)	57.1% male	Rai 0: 77.8%; Rai I-IV: NR	NR	prospective cohort	median 42 months (range 1 - 82 months; 2038 person-years)	
	Narrative	Pflug 2014 (Mayo cohort)	NR	USA	190 events 676 persons	median 61.5 years (range: 32 - 89 years)	67% male	Rai 0: 57.1% Rai I: 34% Rai II: 5.9% Rai III: 1% Rai VI: 1.9%	NR.	prospective cohort	median 57 months (range not reported)	
		Rani 2018 (Indian cohort)	NR	India	86 events 198 persons	median 60 years (range: not reported) hamide-rituximab; CHL:	77% male	Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%	62 CHL 56 rituximab-based therapy 20 other therapies	NR	median 40.5 months (range 1 - 215 months)	

Rossi (Rossi 2013) (Rossi D - Rossi 2013 (Italian cohort))

The Rossi model predicts OS and was derived from data from a retrospective, multicentre cohort study conducted in Italy (Rossi D-Rossi 2013 (Italian cohort)). A total of 637 individuals were included for the model development. The recruitment period for the study lasted from 1996 to 2011. The median follow-up time was 67.2 months; ranges were not reported. All disease stages were included; most individuals were classified as Rai 0 stage (74.9%). The final model included five predictors: TP53, BIRC3 DIS, SF3B1 M, NOTCH1

M, and del11q22-q23. We identified two external validations (Rossi V - Jeromin 2014 (Munich cohort); Rossi V - Rossi 2013 (unclear)). One validation study included a total of 1160 individuals presenting between 2005 and 2010 (Rossi V - Jeromin 2014 (Munich cohort)). The second validation study was conducted between 1996 and 2011; the authors failed to report the number of participants included in the validation cohort (Rossi V - Rossi 2013 (unclear)). We provide a summary of the main characteristics of studies regarding the Rossi model in Figure 6.

Figure 6. Rossi model - summary of characteristics of included studies

Characteristics of	included studies	for the Rossi extern	nally validated r	nodel							
							Characteristi	cs of included stud	ies		
Мо	Model Study		recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up
Rossi model (overall survival)		Rossi 2013 (Development cohort)	1996 - 2011	Italy	178 events 583 persons	51.2% >= 65 years	58.4% male	Rai 0: 74.9% Rai I-II: 11.5%	122 FCR, FR, or pentostatin + CR 64 F or FC 80 alkylator-based (e.g. CHL)	retrospective cohort	median 67.2 months (range not reported)
			2005 - 2010	Unclear	NR events 930 persons	median 67 years (range: 29.6 - 90.5 years)	64.6% male	NR	NR	prospective cohort	median 55.2 months, (range not reported)
		Rossi 2013 (External validation cohort)	1996 - 2011	Italy	62 events 370 persons	NR	55.9% male	Rai 0-1: 75.4% Rai II: 17.3% Rai III-IV: 7.3%	NR	retrospective cohort	median 70.8 months, (range not reported)
Abbreviations: NR: n	ot reported; F: flud	arabine; FC: fludarabin	e + cyclophospha	mide; FCR: fluda	rabine + cyclophosp	hamide + rituximab; FR:	fludarabineritu	ximab, CR: cyclopi	hosphamide + rituximab; CHL: chlora	mbucil	

Stephens model (Stephens 2015) (Stephens OS D - Stephens 2015 (Ohio cohort))

Stephens and colleagues developed models for two outcomes, OS and TFS (Stephens OS D - Stephens 2015 (Ohio cohort); Stephens TFS D - Stephens 2015 (Ohio cohort)). They were derived from data from a retrospective, single-centre cohort study conducted in the USA. A total of 114 individuals were included for the model development. The recruitment period for the study lasted from 2002 to 2012. The follow-up time was not reported. All disease stages were included; most individuals were classified as Rai I/II

stage (46%). The final models for both outcomes were simplified for use in clinical practice. The simplified risk score for OS includes three predictors: ECOG PS, age, and lactate dehydrogenase. The simplified risk score for TFS included five predictors: ECOG PS, Rai stage, age, white blood cell count, and del11q22.3. We identified one external validation study (Stephens OS V - Stephens 2015 (MDACC)), which included 129 individuals. The recruitment period of the validation study was not reported. We provide a summary of the main characteristics of studies regarding the Stephens model in Figure 7.



Figure 7. Stephens model for OS - summary of characteristics of included studies

Characteristics of	Characteristics of included studies for the Stephens externally validated model											
							Characteristi	cs of included stud	ies			
Мо	del	Study	recruitment	ment country number of events age gender stage treatment design								
Stephens model (overall survival)		Stephens 2015 (Development cohort)	2002 - 2012	USA		median 62 years (range: 40 - 92 years)	1	II: 46%; Rai III-	35 purine analogs 18 treatment on a clinical trial 5 other treatment	retrospective cohort	NR	
		Stephens 2015 (MDACC cohort)	NR	USA		median 63 years (range: 40 - 85 years)		Rai 0: 29%; Rai I- II: 52%; Rai III- IV: 19%	INR	retrospective cohort	NR	
Abbreviations: NR: r	breviations: NR not reported											

Prognostic models for the prediction of PFS or TFS (without metaanalysis, 1 to 3 external validations)

Baliakas model (Baliakas 2019) (Baliakas D - Baliakas 2019 (multicentre))

The Baliakas model predicts TFS and was derived from data from a retrospective, multicentre cohort study conducted in 10 European institutions (Baliakas D - Baliakas 2019 (multicentre)). A total of 1900 individuals were included for the development of two separate models, dividing the study population into mutated (M-CLL) and unmutated CLL (U-CLL) patients. The recruitment period for the study was not reported. The median follow-up time

was 7.1 years, ranging from 0.1 to 33.1 years. Only individuals with disease stage Binet A were included. The final model for M-CLL included three predictors: TP53 abnormality, trisomy 12, and stereotyped subset #2 defined as IGHV3-21/IGLV3-21 BcR IG. The final model for U-CLL included three predictors: TP53 abnormality, del11q, and gender. We identified one external validation study (Baliakas V - Baliakas 2019 (MLL + Scan.)), which included a total of 649 persons from two separate studies, one conducted at the Munich Leukemia Laboratory (508 persons) and one Scandinavian population-based study (141 persons). We provide a summary of the main characteristics of studies regarding the Baliakas model in Figure 8.

Figure 8. Baliakas model - summary of characteristics of included studies

Characteristics of	Characteristics of included studies for the Baliakas externally validated model											
				Characteristics of included studies								
Mo	del	Study	recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up	
Baliakas model		Baliakas 2019 (Development cohort)	NR	multiple European countries	NR events 1900 persons	median 64.3 years (range: 22 - 92 years)	61% male	Binet A: 80% Binet B: 12% Binet C: 8%	NR	retrospective cohort	median 7.1 years (range 0.1 - 33.1 years)	
(time-to-first- treatment)		Baliakas 2019 (MLL + Scan. cohort)	NR.			median 63.6 years (range: 29 - 89 years)	62% male	Binet A: 100%	NR	retrospective cohort	NR	
Abbreviations: NR:	breviations: NR: not reported											

GIMEMA model (Molica 2005) (GIMEMA D - Molica 2005 (GIMEMA cohort))

The GIMEMA model predicts PFS and was derived from data from a retrospective, multicentre cohort study conducted in Italy (GIMEMA D - Molica 2005 (GIMEMA cohort)). A total of 1138 individuals were included for the model development. The recruitment period for the study lasted from 1991 to 2000. The median follow-up time was 54 months, ranging from 4 to 309 months. Only individuals with

disease stage Binet A were included. The final model included four predictors: lymphocyte doubling time, absolute peripheral blood lymphocytosis, Rai stage, and gender. We identified one external validation study (GIMEMA V - González Rodríguez 2009 (Cabueñes coh.)), which included 265 persons presenting between 1997 and 2007. We provide a summary of the main characteristics of studies regarding the GIMEMA model in Figure 9.

Figure 9. GIMEMA model - summary of characteristics of included studies

Characteristics of	included studies	for the GIMEMA ex	ternally validat	ted model								
				Characteristics of included studies								
Мо	Model Study		recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up	
GIMEMA model (progression-free survival)		Molica 2005 (Development cohort)	1991 - 2000	litaly		median 65 years (range: 27 - 100 years)	54% male	Rai 0: 77.2 % Rai I-III: 22.7%	133 CHL 20 F or FC 7 CHOP-like regimen 40 other (CVP, C+/-prednisone)	retrospective cohort	median 54 months, range 4 - 309 months	
		González Rodríguez 2009 (Cabueñes hospital cohort)	1997 - 2007	Spain		median 71.7 years (range: 42 - 94 years)	NR	Binet A: 76.8% Binet B: 14.8% Binet C: 8.4%	NR	retrospective cohort	NR	
Abbreviations: NR: r	ot reported; CHL: ch	nlorambucil; F: fludarat	oine; FC: fludarabi	ine + cyclophosp	hamide; CHOP: dox	orubicin + cyclophospha	mide + vincristi	ne + prednisone; (OVP: cyclophosphamide + vincristine :	sulfate + predniso	ne	

MDACC 2011 (Wierda 2011) (MDACC 2011 D - Wierda 2011 (MDACC))

The MDACC 2011 model predicts TFS and was derived from data from a retrospective, single-centre cohort study conducted in the USA (MDACC 2011 D - Wierda 2011 (MDACC)). A total of 930 individuals were included for the model development. The recruitment period for the study lasted from 2004 to 2009. The median follow-up time was 26 months, ranging from three to

73 months. All disease stages were included; most individuals were classified as Rai I stage (51%). The final model included six predictors: IgHV mutational status, diameter of largest palpated lymph node, FISH category (del11q or del17p versus none), number of involved lymph node sites, lactate dehydrogenase (LDH), and the IgHV mutational status and LDH interaction term. We identified one external validation study (MDACC 2011 V - Molica 2016 (O-CLL1-



GISL)), which included 328 persons presenting between 2006 and

2010. We provide a summary of the main characteristics of studies regarding the MDACC 2011 model in Figure 10.

Figure 10. MDACC 2011 model - summary of characteristics of included studies

Characteristics of	haracteristics of included studies for the MDACC 2011 externally validated model											
				Characteristics of included studies								
Мо	del	Study	recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up	
MDACC 2011 model (time-to-first- treatment)	1	Wierda 2011 (Development cohort)	2004 - 2009	USA		median 59 years (range: 30 - 89 years)	61% male	Rai 0: 36% Rai I: 51% Rai II: 8% Rai III-IV: 5%	NR	retrospective cohort	median 26 months (range 3 - 73 months)	
		Molica 2016 (O-CLL1- GISL)	2006 - 2010	Italy		median 61 years (range: 33 - 70 years)	58.2% male	Rai 0: 76.5% Rai I-II: 23.5%	NR	prospective cohort	median 30 months (range 1 - 65 months)	
Abbreviations: NR: r	reviations: NR: not reported											

Morabito model (Morabito 2009) (Morabito D - Morabito 2009 (Italian cohort))

The Morabito model predicts TFS and was derived from data from a multicentre study conducted in Italy (Morabito D - Morabito 2009 (Italian cohort)). The study design was not reported. A total of 262 individuals were included for the model development. The recruitment period was not reported. The median follow-up time was 36 months, ranging from 12 to 180 months. Only individuals

with disease stage Binet A were included. The final model included three predictors: IgHV mutational status, CD38, and ZAP-70. We identified one external validation study (Morabito V - Gentile 2014 (O-CLL1-GISL)), which included 480 persons presenting from 2007. There was no information regarding the end of the recruitment period. We provide a summary of the main characteristics of studies regarding the Morabito model in Figure 11.

Figure 11. Morabito model - summary of characteristics of included studies

Characteristics of	characteristics of included studies for the Morabito externally validated model										
				Characteristics of included studies							
Мо	del	Study	recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up
Morabito model (time-to-first- treatment)	1	Morabito 2009 (Development cohort)	NR	Italy		median 65 years (range not reported)	61.5% male	Binet A: 100%	NR	NR	median 36 months (range 12 - 180 months)
		Gentile 2014 (O-CLL1- GISL)	2007 - NR	Italy	121 events 468 persons	47.4 % < 60 years	58.5% male	Rai 0: 77.3% Rai I-II: 22.7%	NR.	prospective cohort	median 38.5 months (range 6 - 82 months)
Abbreviations: NR: r	verviations: NR: not reported										

O-CLL1 model (Gentile 2016) (O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL))

The O-CLL1 model predicts TFS and was derived from data from a prospective, multicentre cohort study conducted in Italy (O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL)). A total of 480 individuals were included for the model development. Recruitment for the study started in 2007; the end date was not reported. The median follow-up time was 42 months, ranging from 6 to 82 months. Only individuals with disease stage Binet A were included. The final model included four predictors: Rai stage, absolute lymphocyte

count (ALC), serum B2-microglobulin, and IgHV mutational status. We identified two external validation cohorts for TFS (O-CLL1 V - Gentile 2016 (Mayo cohort); O-CLL-1 V - Rani 2018 (Indian cohort)), which included a total of 626 persons. One validation study recruited participants between 2001 and 2008 (O-CLL1 V - Gentile 2016 (Mayo cohort)); the other study lacked information regarding the recruitment period (O-CLL-1 V - Rani 2018 (Indian cohort)). We provide a summary of the main characteristics of studies regarding the O-CLL1 model in Figure 12.

Figure 12. O-CLL-1 model - summary of characteristics of included studies

Characteristics of	included studies	for the O-CLL1 exte	rnally validated	d model							
							Characteristi	cs of included stud	ies		
Mod	del	Study	Study recruitment		number of events / number	age	gender	stage	treatment	design	follow-up
O-CLL1 model (time-to-first-		Gentile 2016 (O-CLL1- GISL)	2007 - NR	Italy	84 events 337 persons	46.7% <= 60 years	57.3% male	Rai 0: 77.7% Rai I-II: 22.3%	NR	prospective cohort	median 42 months (range 6 - 82 months)
treatment)		Rani 2018 (Indian cohort)	NR	India	86 events 198 persons	median 60 years (range: not reported)	77% male	34.9%; Rai III:	62 CHL	NR	median 40.5 months (range 1 - 215 months)
		Gentile 2016 (Mayo cohort 2001-2008)	2001 - 2008	USA	130 events 428 persons	40.2% <= 60 years	66.4% male	Rai 0: 67.5% Rai I-II: 32.5%	NR	NR	median 97 months (range not reported)
Abbreviations: NR: n	ot reported; CHL: d	hlorambucil							·		

Stephens model (Stephens 2015) (Stephens TFS D - Stephens 2015 (Ohio cohort))

For a description of the model for TFS in Stephens and colleagues (Stephens TFS D - Stephens 2015 (Ohio cohort)), please see

description in section OS (above), as the authors used the same development and validation cohorts for both outcomes. We provide a summary of the main characteristics of studies regarding the Stephens model in Figure 13.



Figure 13. Stephens model for TFS - summary of characteristics of included studies

Characteristics of	Characteristics of included studies for the Stephens externally validated model										
							Characteristi	cs of included stud	ies		
Мо	del	Study	recruitment	country	number of events / number	age	gender	stage	treatment	design	follow-up
Stephens model (treatment-free survival)		Stephens 2015 (Development cohort)	2002 - 2012	USA	58 events 114 persons	median 62 years (range: 40 - 92 years)		II: 46%; Rai III-	35 purine analogs 18 treatment on a clinical trial 5 other treatment	retrospective cohort	NR
	1	Stephens 2015 (MDACC cohort)	NR	USA	NR events 129 persons	median 63 years (range: 40 - 85 years)	1	Rai 0: 29%; Rai I- II: 52%; Rai III- IV: 19%	NR	retrospective cohort	NR
Abbreviations: NR: r	ot reported										

Models without external validation studies

The 34 references with 41 model or score developments can be found in Studies awaiting classification, as it is widely recommended that developed prediction models should not be used in daily practice before they were validated at least once (Moons 2009; Moons 2015; Moons 2019; Steyerberg 2013). We have provided an overview of the main characteristics of these developed (but never validated) models in Appendix 7. In total, 19 models were developed to predict OS, 19 to predict TFS and three to predict PFS. These models were published between 1982 and 2019 and contained an average number of 3.8 predictors (range between 2 and 6). Nineteen models were derived from retrospective cohorts, three were on prospective cohorts and 19 did not report the study design clearly. The most commonly included predictors were IgHV status (mutated versus unmutated), B2-microglobulin, age, clinical stage, genomic aberrations as defined by Döhner 2000, ZAP-70 expression, CD38 expression and gender.

Excluded studies

During abstract screening, we excluded 18,713 references that clearly did not match our inclusion criteria. Of the remaining 283 full-text references, we excluded 213 for the following reasons.

References of prognostic factor studies or prognostic factor identification studies (130 references) (Berke 2019; Bo 2014; Brejcha 2010; Brugiatelli 2007; Bulian 2014; Byrd 2006; Cailliod 2005; Callea 1999; Catovsky 1989; Cesano 2013; Chang 2003; Chauzeix 2018; Chen 1997; Chena 2008; Chevallier 2002; Chiaretti 2014; Christiansen 1994; Ciccone 2012; Claus 2012; Claus 2014; Cmunt 2002; Cocco 2005; Corcoran 2005; Cordone 1998; Cortese 2014; Coscia 2012; Crespo 2003; Cro 2009; D'Arena 2001; D'Arena 2007; Damle 1999; DeAndres-Galiana 2016; Degan 2004; Delgado 2009; Delgado 2014; Del Guidice 2011; Del Poeta 2010; Del Principe 2004; Del Principe 2006; Di Raimondo 2001; Dong 2011; Dong 2014; Durak 2009; El-Kinawy 2012; Gattei 2008; Gdynia 2018; Gentile 2016; Giudice 2018; Gogia 2014; Grabowski 2005; Han 1984; Hock 2010; Hus 2006; Jaksic 1981; Josefsson 2007; Juliusson 1986; Juliusson 1990; Kahraman 2014; Kardum-Skelin 2008; Karmiris 1994; Khalifa 2002; Kim 2004; Kimby 1988; Knospe 1977; Koberda 1989; Korycka-Wolowiec 2011; Krober 2002; Kryachok 2011; Kurec 1992; Lai 2002; Lech-Maranda 2012; Lech-Maranda 2013; Lecouvet 1997; Li 2008; Li 2017a; Lin 2002; Lin 2014; Lozano-Santos 2014; Lucas 2015; Maffei 2007; Maffei 2010; Mansouri 2013; Marasca 2005; Marasca 2013; Martinelli 2008; Masic 1998; Mateva 2001; Matthews 2006; Matthews 2007; Matutes 2013; Miao 2018; Molica 1986; Molica 1988; Molica 1991; Molica 1994; Molica 1998; Molica 1999a; Molica 1999b; Molica 2008; Montserrat 1991; Morabito 2001; Morabito 2015a; Morabito 2018c; Morilla 2008; Nenova 2000; Nipp 2014; Nowakowski 2009; Nuckel 2006; Nuckel 2009; Ocana 2007; Oliveira 2011; Oscier 1990; Paolino 1984; Prokocimer 1985; Qin 2017; Resegotti 1989; Rissiek 2014; Ronchetti 2016; Sarmiento 2002; Shanafelt 2010; Spacek 2009; Stamatopoulos 2017; Strefford 2015; Szymczyk 2018; Vojdeman 2017; Vural 2014; Weiss 2011; Wierda 2003; Winkler 2010; Zenz 2009), and five references aimed at identifying prognostic factor thresholds (Dasgupta 2015; Davis 2016; Rossi 2010a; Tobin 2005a; Tobin 2005b).

- The main focus was on the development of a diagnostic CLL staging system (21 references) (Apelgren 2006; Baccarani 1982; Binet 1981; Binet 1977; Chelazzi 1979; Ciocoiu 1988; De Faria 2000; De Rossi 1989; Ferrara 1981; Jaksic 1992; Molica 1984; Moreno 2019; Rai 1990; Rai 1975; Rossi 1986; Rozman 1979; Santoro 1979; Scolozzi 1981; Velardi 1980; Wu 2010; Zengin 1997).
- Involved genetic analysis only (16 references), e.g. genetic subgrouping, genetic signature(s) or genetic clustering (Baliakas 2015; Bomben 2009; Bou Samra 2014; Chuang 2012; Ferreira 2014; Friedman 2009; Herold 2011; Houldsworth 2014; Morabito 2015b; Orgueira 2019; Queiros 2015; Raponi 2018; Rodriguez 2007; Vallat 2013; Van Damme 2012; Zucchetto 2006).
- Included a population which did not match our PICOTS question (10 references), e.g. previously treated individuals with CLL (Giles 2003; Kardum-Skelin 2009; Keating 2000; Krober 2006; Melo 1987; Nola 2004; O'Brien 1993; Rossi 2010b; Rossi 2011; Weinberg 2007).
- Nine references focused on the characterisation of CLL and the prevalence of prognostic factors (Criel 1997; Cro 2010; Cuneo 2004; D'Arena 2012; Geisler 1997; Gonzalez 2013; Gonzalez-Gascon 2015; Gonzalez-Rodriguez 2010; Hallek 1999).
- Nine references of other types of studies, e.g. risk factor study, diagnostic accuracy study (Cuneo 2018; Deslandes 2007; Dimier 2018; Fang 2019; Kay 2018; Kleinstern 2018; Plesingerova 2017; Salomon-Nguyen 1995; Savvopoulos 2016).
- Four references developed or used an outdated staging system, which is no longer used in today's clinical practice (Bettini 1986; Chastang 1985; French Cooperative Group on CLL 1988; Mandelli 1987).
- Five other types of references, e.g. review or comment to an excluded study (Bomben 2005; Grever 2006; Jaksic 2014; Matutes 2017; Nedeva 2018).
- Three references evaluated outcomes not relevant to this review (Shanafelt 2017; Stamatopoulos 2009; Tallarico 2018).
- Two references focused on scores or models which included predictors available at treatment initiation only (incorrect timing) (Gentile 2018; Nabhan 2017).



Reporting deficiencies

Appendix 8 describes the reporting deficiencies of the included studies per model for the primary outcome each model was developed for. The number of studies includes the development study. Data obtained by contacting authors are included as available.

For the Barcelona-Brno model, the number of events for two studies and calibration for one study was obtained by contacting the corresponding authors of the primary studies. For the CLL-IPI, information on total sample size and number of events was provided by one author. Information on calibration was not reported in publications at all, but was obtained by contacting the corresponding authors. For the GCLLSG model, the confidence intervals of the c-statistic and calibration for the development study publication with one external validation was obtained from the authors. For the MDACC 2007 model, the c-statistic and 95% CI for one external validation study was provided by the corresponding author. We did not obtain any additional information for the other models. We received primary data of one study, but were unable to reconstruct the analysis.

Of the 41 models without external validations, all studies reported the total number of individuals included for analysis. Eighteen studies lacked information on the recruitment period, 19 did not clearly report their study design, 24 did not report the calibration of their model and 31 did not report any measure of discrimination for their model (Appendix 7). Since we included studies with a wide range of publication years, the large amount of missing information may be a result of the methodological evolution of prognostic model research. Moreover, with our decision to also include scores where authors decided seemingly ad hoc to assign points and create a prognostic score, we have included studies that show more resemblance with prognostic factor studies (i.e. testing of the independence of factors from each other) than with a prognostic model development.

Summarised over all categories of included studies (development studies without external validations (N = 40), development studies with external validations (N = 12), external validations of primary

outcome of an included model (N = 35)), information was especially lacking on calibration and discrimination. Calibration was reported in 17 and discrimination in 45 out of 87 studies.

Risk of bias and applicability assessment of included studies

Models with more than three external validation studies per outcome

Prognostic models for the prediction of OS (with meta-analysis, > 3 external validations)

CLL-IPI (CLL-IPI D - Bahlo 2016 (development cohort))

We rated the risk of bias of the development study as low for the domains: participants, predictors and outcome (CLL-IPI D - Bahlo 2016 (development cohort)), and high for the domain 'analysis' due to several reasons (univariable selection and dichotomisation of factors, missing data handling). We rated the risk of bias for the validation studies as low for the domain 'participants' for two studies (CLL-IPI V - Gentile 2016 (Italian cohort); CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort)), unclear in the case of six studies due to unclear eligibility criteria, and high for two studies due to inappropriate inclusion criteria (CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014); CLL-IPI V - Rani 2018 (Indian cohort)). We rated the risk of bias for the domains 'predictors' and 'outcome' as low for most studies, except one study with an unclear rating for the domain 'predictors' because predictor assessment has probably changed (CLL-IPI V - Gentile 2016 (Italian cohort)), and three studies with an unclear rating for the domain 'outcome' due to short observation time (CLL-IPI V - Da Cunha-Bang 2016 (Danish cohort); CLL-IPI V -Rani 2018 (Indian cohort); CLL-IPI V - Rigolin 2017 (Ferrera cohort)). Concerning the domain 'analysis', we considered seven studies to have a high risk of bias due to inappropriate handling of missing data, low number of events and lack of reporting of performance measures. Three studies had a low rating (CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014); CLL-IPI V - Bahlo 2016 (SCAN cohort); CLL-IPI V - Delgado 2017 (Barcelona cohort)), and one had an unclear rating due to lack of performance measures (CLL-IPI V - Da Cunha-Bang 2016 (Danish cohort)). One validation study did not examine the outcome of interest (CLL-IPI V - Molica 2016 (O-CLL1-GISL)) (Figure 14).

Figure 14. Risk of bias (PROBAST) assessment of the CLL-IPI model (Bahlo 2016)

			Pre	ediction model Ris	k Of Bias ASsessn	nent Tool (PROBA	ST)
Mode	d .	Study	Domain 1: Participants	Domain 2: Predictors	Domain 3: Outcome	Domain 4: Analysis	Overall judgement
CLL-IPI model	Development	Bahlo 2016 (Development cohort)	low	low	low	high	high
(overall survival)		Bahlo 2016 (Mayo clinic 2001-2014)	high	low	low	low	high
		Bahlo 2016 (SCAN cohort)	unclear	low	low	low	unclear
	Meta-	Delgado 2017 (Barcelona cohort)	unclear	low	low	low	high
	analysed	Gentile 2016 (Italian cohort)	low	unclear	low	high	high
	o-statistic	Muñoz-Novaz 2018 (Spanish cohort)	low	low	low	high	high
		Rani 2018 (Indian cohort)	high	low	unclear	high	high
		Zhu 2018 (Chinese cohort)	unclear	low	low	high	high
		Da Cunha-Bang 2016 (Danish cohort)	unclear	low	unclear	unclear	unclear
	Narrative	Molica 2016 (O-CLL1-GISL)	NA	NA	NA	NA	NA
	Ivarrative	Reda 2017 (Milan cohort)	unclear	low	low	high	high
		Rigolin 2017 (Ferrera cohort)	unclear	low	unclear	high	high

We rated the concern for applicability of the development study as high for the domain 'participants' due to the high proportion of individuals with treatment indication (CLL-IPI D - Bahlo 2016 (development cohort)). We rated this low for the domains 'predictors' and 'outcome'. We rated concern for applicability as unclear for the domain 'participants' in all validation studies but two, which received a low rating (CLL-IPI V - Da Cunha-Bang 2016

(Danish cohort); CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort)), due to unclear eligibility criteria. We rated concern for applicability as unclear for the domain 'predictors' in six validation studies because one predictor (TP53 mutation) was replaced by a proxy predictor (del(17p). The remaining four were rated as low (CLL-IPI V - Bahlo 2016 (SCAN cohort); CLL-IPI V - Delgado 2017 (Barcelona cohort); CLL-IPI V - Rigolin 2017 (Ferrera cohort); CLL-IPI V - Zhu



2018 (Chinese cohort)). We considered the concern for applicability to be low for the domain 'outcome' in all validation studies. One

validation study (CLL-IPI V - Molica 2016 (O-CLL1-GISL))) did not examine the outcome of interest (Figure 15).

Figure 15. Applicability assessment for all developed models with external validations

			PR	DBAST - Applicab	ility
Model		Study	Domain 1: Participants	Domain 2: Predictors	Domain 3: Outcome
Baliakas model	Development	•	high	low	low
(time-to-first-treatment)	Narrative	Baliakas 2019 (MLL + Scan. cohort)	unclear	low	low
Barcelona-Brno model	Development	Delgado 2017 (Development cohort)	unclear	low	low
overall survival)	Meta-	Delgado 2017 (Brno cohort, CZ)	unclear	low	low
	analysed	Gentile 2017 (Italian multicentre and Mayo cohort)	unclear	low	low
	c-statistic	Muñoz-Novaz 2018 (Spain multicentre cohort)	low	low	low
		Rani 2018 (Indian cohort)	unclear	unclear	low
	Narrative	Molica 2017 (O-CLL1-GISL)	NA	NA	high
		Reda 2017 (Milan cohort)	unclear	unclear	low
CLL-IPI model	Development	Bahlo 2016 (Development cohort)	high	low	low
overall survival)		Bahlo 2016 (Mayo clinic 2001-2014)	unclear	unclear	low
		Bahlo 2016 (SCAN cohort)	unclear	low	low
	Meta-	Delgado 2017 (Barcelona hospital cohort)	unclear	low	low
	analysed	Gentile 2016 (Italian multicentre cohort)	unclear	unclear	low
	c-statistic	Muñoz-Novaz 2018 (Spain multicentre cohort)	low	unclear	low
		Rani 2018 (Indian cohort)	unclear	unclear	low
		Zhu 2018 (Chinese cohort)	unclear	low	low
		Da Cunha-Bang 2016 (Danish cohort)	low	unclear	low
	Narrative	Molica 2016 (O-CLL1-GISL)	NA	NA	high
	Traina.ire	Reda 2017 (Milan cohort)	unclear	unclear	low
		Rigolin 2017 (Ferrera cohort)	unclear	low	low
ADACC 2007 model	Development	Wierda 2007 (Development cohort)	low	low	low
overall survival)	1	Bulian (Italian-Swiss cohort)	low	low	low
	Meta- analysed	Gentile 2014 (Italian multicentre cohort)	unclear	low	low
	c-statistic for	Gentile 2016 (Mayo cohort 2001-2008)	unclear	low	low
	the index	González Rodríguez 2009 (Cabueñes hospital cohort)	low	low	low
	score	Muñoz-Novaz 2018 (Spain multicentre cohort)	low	low	low
		Rani 2018 (Indian cohort)	unclear	unclear	low
		Molica 2010 (GIMEMA cohort 1991-2000)	low	high	high
	Narrative	Molica 2015 (O-CLL1-GISL)	NA	NA NA	high
	Trainadire	Pflug 2014 (German CLL study group cohort)	unclear	low	low
		Trajkova 2013 (Macedonian cohort)	low	low	low
GCLLSG model	Development	Pflug 2014 (Development cohort)	unclear	low	low
overall survival)		Molica 2015 (O-CLL1-GISL)	NA	NA NA	high
	Narrative	Pflug 2014 (Mayo cohort)	unclear	low	low
		Rani 2018 (Indian cohort)	NA	NA	high
GIMEMA model	Development	Molica 2005 (Development cohort)	low	low	low
progression-free survival)	Narrative	González Rodríguez 2009 (Cabueñes hospital cohort)	low	low	low
MDACC 2011 model	Development	Wierda 2011 (Development cohort)	unclear	low	low
time-to-first-treatment)	Narrative	Molica 2013 (O-CLL1-GISL)	low	low	low
Norabito model	Development	Morabito 2009 (Development cohort)	unclear	high	low
time-to-first-treatment)	Narrative	Gentile 2014 (O-CLL1-GISL)	low	low	low
D-CLL1model	Development	Gentile 2016 (O-CLL1-GISL)	low	low	low
time-to-first-treatment)		Rani 2018 (Indian cohort)	unclear	unclear	low
	Narrative	Gentile 2016 (Mayo cohort 2001–2008)	unclear	low	low
Rossi model	Development	Rossi 2013 (Development cohort)	low	low	low
overall survival)		Jeromin 2014 (Munich cohort)	low	low	low
	Narrative	Rossi 2013 (External validation cohort)	low	low	low
Stephens model	Development	Stephens 2015 (Development cohort)	high	low	low
overall survival)	Narrative	Stephens 2015 (MDACC cohort)	high	low	low
Stephens model	Development	Stephens 2015 (NDACC condit)			
treatment-free survival)	Narrative		high bigh	low	low
	ivariative	Stephens 2015 (MDACC cohort)	high	low	low

Barcelona-Brno model (Barcelona-Brno D - Delgado 2017 (Barcelona cohort))

We rated the risk of bias of the development study as unclear for the domain 'participants' due to missing information on eligibility criteria (Barcelona-Brno D - Delgado 2017 (Barcelona cohort)), low for the domains 'predictors' and 'outcome', and high for the domain

'analysis' due to the model-building process (factor selection and weighting). We rated the risk of bias between the validation studies as unclear for the domain 'participants' in three studies (Barcelona-Brno V - Delgado 2017 (Brno cohort); Barcelona-Brno V - Rani 2018 (Indian cohort); Barcelona-Brno V - Reda 2017 (Milan cohort)) due to missing eligibility criteria and recruitment period, and low



in two studies (Barcelona-Brno V - Gentile 2017 (Italian & Mayo); Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.)). We rated the risk of bias for the domains 'predictors' and 'outcome' as low for all validation studies except one (Barcelona-Brno V - Rani 2018 (Indian cohort)), which we rated as unclear for the domain 'outcome' due to the short observation time. Concerning the domain 'analysis',

we considered all studies to be at high risk of bias (especially due to handling missing data, low number of events and reporting of performance measures) except one (Barcelona-Brno V - Delgado 2017 (Brno cohort)), which we rated as unclear. One validation study did not examine the outcome of interest (Barcelona-Brno V - Molica 2017 (O-CLL1-GISL)) (Figure 16).

Figure 16. Risk of bias (PROBAST) assessment of the Barcelona-Brno model (Delgado 2017)

			Pre	ediction model Ris	k Of Bias ASsessn	ent Tool (PROBA	ST)
Model		Study	Domain 1: Participants	Domain 2: Predictors	Domain 3: Outcome	Domain 4: Analysis	Overall judgement
Barcelona-Brno model	Development	Delgado 2017 (Development cohort)	unclear	low	low	high	high
(overall survival)	м.	Delgado 2017 (Brno cohort, CZ)	unclear	low	low	unclear	unclear
	Meta- analysed	Gentile 2017 (Italian multicentre and Mayo cohort)	low	low	low	high	high
	c-statistic	Muñoz-Novaz 2018 (Spanish cohort)	low	low	low	high	high
	O Statistic	Rani 2018 (Indian cohort)	unclear	low	unclear	high	high
	Narrative	Molica 2017 (O-CLL1-GISL)	NA	NA	NA	NA	NA
	Ivanadve	Reda 2017 (Milan cohort)	unclear	low	low	high	high

We rated the concern for applicability of the development study as unclear for the domain 'participants' and low for the domains 'predictors' and 'outcome' (Barcelona-Brno D - Delgado 2017 (Barcelona cohort)). Similarly, we rated concern for applicability as unclear for the domain 'participants', and low for the domains 'predictors' and 'outcome' in three validation studies (Barcelona-Brno V - Delgado 2017 (Brno cohort); Barcelona-Brno V - Gentile 2017 (Italian & Mayo); Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.)) due to undefined eligibility criteria. We assigned two validation studies (Barcelona-Brno V - Rani 2018 (Indian cohort); Barcelona-Brno V - Reda 2017 (Milan cohort)) the rating unclear for the domain 'predictors' due to lack of information on the timing of predictor assessment. One validation study did not examine the outcome of interest (Barcelona-Brno V - Molica 2017 (O-CLL1-GISL)) (Figure 15).

MDACC 2007 (Wierda 2007) (MDACC 2007 D - Wierda 2007 (MDACC))

We rated the risk of bias for the development study as low across all domains (MDACC 2007 D - Wierda 2007 (MDACC)). We rated the

risk of bias across the validation studies as low for the domain 'participants' in six validation studies and high in three validation studies (MDACC 2007 V - Gentile 2016 (Mayo cohort); MDACC 2007 V - Rani 2018 (Indian cohort); MDACC 2007 V - Trajkova 2013 (Macedonia)) due to unclear eligibility criteria. We considered all validation studies to have a low risk of bias for the domain 'predictors'. We rated three studies as unclear for the domain 'outcome' (MDACC 2007 V - Molica 2010 (GIMEMA cohort); MDACC 2007 V - Rani 2018 (Indian cohort); MDACC 2007 V - Trajkova 2013 (Macedonia)) because of the short observation time. Concerning the domain 'analysis', we considered all validation studies to have a high or unclear risk of bias due to the low number of events and inappropriate handling of missing data. One validation study did not examine the outcome of interest (MDACC 2007 V - Molica 2015 (O-CLL1-GISL)) (Figure 17).

Figure 17. Risk of bias (PROBAST) assessment of the MDACC 2007 model (Wierda 2007)

			Pre	ediction model Ris	k Of Bias ASsessn	nent Tool (PROBA:	ST)
Model		Study	Domain 1: Participants	Domain 2: Predictors	Domain 3: Outcome	Domain 4: Analysis	Overall judgement
	Development	Wierda 2007 (Development cohort)	low	low	low	low	low
(overall survival)		Bulian (Italian-Swiss cohort)	low	low	low	high	high
	Meta-	Gentile 2014 (Italian multicentre cohort)	low	low	low	high	high
		Gentile 2016 (Mayo cohort 2001-2008)	high	low	low	unclear	high
		González Rodríguez 2009 (Cabueñes hospital cohort)	low	low	low	high	high
	the index	Muñoz-Novaz 2018 (Spain multicentre cohort)	low	low	low	high	high
	score	Pflug 2014 (German CLL study group cohort)	low	low	low	high	high
		Rani 2018 (Indian cohort)	high	low	unclear	high	high
		Molica 2010 (GIMEMA cohort 1991-2000)	low	low	unclear	high	high
	Narrative	Molica 2015 (O-CLL1-GISL)	NA	NA	NA	NA	NA
		Trajkova 2013 (Macedonian cohort)	high	low	unclear	high	high

We rated the concern for applicability of the development study as low across all domains (MDACC 2007 D - Wierda 2007 (MDACC)). We rated concern for applicability as low for the domain 'participants' in five validation studies and unclear in four validation studies due to unclear eligibility criteria (MDACC 2007 V - Gentile 2014 (Italian cohort); MDACC 2007 V - Gentile 2016 (Mayo cohort); MDACC 2007 V - Pflug 2014 (3 RCTs); MDACC 2007 V - Rani 2018 (Indian cohort)). Concerning the domain 'predictors', we considered the concern for applicability to be low in seven validation studies, high in one study (MDACC 2007 V - Molica 2010 (GIMEMA cohort)), and unclear in one study (MDACC 2007 V - Rani 2018 (Indian cohort)). Concern

for applicability was rated as low for the domain 'outcome' in all validation studies except one, for which we assigned a high rating (MDACC 2007 V - Molica 2010 (GIMEMA cohort)). One validation study did not examine the outcome of interest (MDACC 2007 V - Molica 2015 (O-CLL1-GISL)) (Figure 15).

Prognostic models for the prediction of PFS or TFS (with metaanalysis, > 3 external validations)

We did not identify any prognostic model development studies that were externally validated more than three times for the outcomes PFS, TFS or TFS.



Models with one to three external validation studies per outcome

Prognostic models for the prediction of OS (without meta-analysis, 1 to 3 external validations)

GCLLSG model (Pflug 2014) (GCLLSG D - Pflug 2014 (GCLLSG))

We rated the risk of bias for the development study as low for all domains except 'analysis' (GCLLSG D - Pflug 2014 (GCLLSG)). We rated 'analysis' at high risk of bias due to dichotomisation and

univariable selection of factors, inappropriate handling of missing data and simplification of the model (loss of original weighting). We considered the only validation study with a matching outcome to have a high risk of bias for the domains 'participants' and 'analysis' due to inclusion of participants with available data and low number of events (GCLLSG V - Pflug 2014 (Mayo cohort)). Risk of bias was low for the domains 'predictors' and 'outcome'. The other two validation studies did not examine the outcome of interest (GCLLSG V - Molica 2015 (O-CLL1-GISL); GCLLSG V - Rani 2018 (Indian cohort)) (Figure 18).

Figure 18. Risk of bias (PROBAST) assessment of other models that were externally validated

Model		Study	Prediction model Risk Of Bias ASsessment Tool (PROBAST)				
			Domain 1: Participants	Domain 2: Predictors	Domain 3: Outcome	Domain 4: Analysis	Overall judgement
Baliakas model (time-to-first-treatment)	Development	Baliakas 2019 (Development cohort)	high	low	low	high	high
	Narrative	Baliakas 2019 (MLL + Scan. cohort)	unclear	unclear	unclear	high	high
GCLLSG model (overall survival)	Development	Pflug 2014 (Development cohort)	low	low	low	high	high
	Narrative	Molica 2015 (O-CLL1-GISL)	NA	NA	NA	NA	NA
		Pflug 2014 (Mayo cohort)	high	low	low	high	high
		Rani 2018 (Indian cohort)	NA	NA	NA	NA NA	NA
GIMEMA model (progression-free survival)	Development	Molica 2005 (Development cohort)	low	low	high	high	high
	Narrative	González Rodríguez 2009 (Cabueñes hospital cohort)	low	low	high	high	high
MDACC 2011 model (time-to-first-treatment)	Development	Wierda 2011 (Development cohort)	high	low	high	high	high
	Narrative	Molica 2013 (O-CLL1-GISL)	low	low	high	high	high
Morabito model (time-to-first-treatment)	Development	Morabito 2009 (Development cohort)	unclear	low	low	high	high
	Narrative	Gentile 2014 (O-CLL1-GISL)	low	low	low	unclear	unclear
O-CLL1model (time-to-first-treatment)	Development	Gentile 2016 (O-CLL1-GISL)	low	low	high	high	high
	INarrative	Rani 2018 (Indian cohort)	unclear	low	high	high	high
		Gentile 2016 (Mayo cohort 2001-2008)	high	low	high	unclear	high
Rossi model (overall survival)	Development	Rossi 2013 (Development cohort)	low	low	low	unclear	unclear
	Narrative	Jeromin 2014 (Munich cohort)	unclear	low	low	high	high
		Rossi 2013 (External validation cohort)	unclear	low	low	high	high
Stephens model (overall survival)	Development	Stephens 2015 (Development cohort)	high	low	low	high	high
	Narrative	Stephens 2015 (MDACC cohort)	high	unclear	low	high	high
Stephens model (treatment-free survival)	Development	Stephens 2015 (Development cohort)	high	low	high	high	high
	Narrative	Stephens 2015 (MDACC cohort)	high	unclear	high	high	high

We rated the concern for applicability of the development study as unclear for the domain 'participants' because the sample was based on RCT data that may not be representative for all individuals with CLL (GCLLSG D - Pflug 2014 (GCLLSG)). Concern was low for the domains 'predictors' and 'outcome'. Similarly, we rated concern for applicability of the validation study as unclear for the domain 'participants' due to eligibility criteria and low for the domains 'predictors' and 'outcome' (GCLLSG V - Pflug 2014 (Mayo cohort)). The other two validation studies did not examine the outcome of interest (GCLLSG V - Molica 2015 (O-CLL1-GISL); GCLLSG V - Rani 2018 (Indian cohort)) (Figure 15).

Rossi (Rossi 2013) (Rossi D - Rossi 2013 (Italian cohort))

We rated the risk of bias of the development study as low across all domains except 'analysis' (Rossi D - Rossi 2013 (Italian cohort)), which we considered to be of unclear risk because handling of missing data handling was not reported. We rated the risk of bias for the two validation studies as unclear due to lack of information on study design for the domain 'participants', low for the domains 'predictors' and 'outcome', and high for the domain 'analysis' due to missing data handling and low number of events (Rossi V - Jeromin 2014 (Munich cohort); Rossi V - Rossi 2013 (unclear)) (Figure 18).

We rated the concern for applicability of the development study as low across all domains (Rossi D - Rossi 2013 (Italian cohort)). We rated concern for applicability of the two validation studies as low for the domains 'participants', 'predictors' and 'outcome' (Rossi V - Jeromin 2014 (Munich cohort); Rossi V - Rossi 2013 (unclear)) (Figure 15).

Stephens model (Stephens 2015) (Stephens OS D - Stephens 2015 (Ohio cohort))

We rated the risk of bias for the development study as high for the domains 'participants' (Stephens OS D - Stephens 2015 (Ohio cohort)), because the study included only individuals with del(17p), which implies that only persons with available FISH assessment could have been included. We considered the domains 'predictors' and 'outcomes' to be of low risk. We rated the domain 'analysis' as high due to small sample size, inappropriate missing data handling and inconsistent reporting of performance measures. In a similar manner, we rated the risk of bias for the validation study as high for the domains 'participants' and 'analysis' due to the same reasons, unclear for the domain 'predictors' and low for the domain 'outcome' (Stephens OS V - Stephens 2015 (MDACC)) (Figure 18).

We rated the concern for applicability of the development study (Stephens OS D - Stephens 2015 (Ohio cohort)) and validation study (Stephens OS V - Stephens 2015 (MDACC)) as high for the domain 'participants' due to the selective sample, and low for the domains 'predictors' and 'outcome' (Figure 15).

Prognostic models for the prediction of PFS or TFS (without metaanalysis, 1 to 3 external validations)

Baliakas model (Baliakas 2019) (Baliakas D - Baliakas 2019 (multicentre))

We rated the risk of bias of the development study as low for the domains 'predictors' and 'outcome', and high for the domains 'participants' since participants with missing data were excluded (Baliakas D - Baliakas 2019 (multicentre)). We rated the



domain 'analysis' as high due to inappropriate handling of missing data, and univariable selection of predictors. Due to reporting deficiencies concerning inclusion criteria, predictor assessment and outcome definition of the validation study (Baliakas V-Baliakas 2019 (MLL + Scan.)), we rated the risk of bias as unclear across all domains except 'analysis', which we considered to be of high risk due to lack of reporting performance measures (Figure 18).

We rated the concern for applicability of the development study as high for the domain 'participants' due to the sample not being representative of all CLL patients and low for the domains 'predictors' and 'outcome' (Baliakas D - Baliakas 2019 (multicentre)). We rated as unclear concern for applicability of the validation study for the domain 'participants' due to lack of information about eligibility criteria and recruitment period (Baliakas V - Baliakas 2019 (MLL + Scan.)), and low for the domains 'predictors' and 'outcome' (Figure 15).

GIMEMA model (Molica 2005) (GIMEMA D - Molica 2005 (GIMEMA cohort))

We rated the risk of bias for the development study as low for the domains 'participants' and 'predictors' (GIMEMA D - Molica 2005 (GIMEMA cohort)), and high for the domains 'outcome' because a predictor was not excluded from the outcome definition and 'analysis' due to inappropriate handling of missing data, univariable selection of predictors, and simplification of the model. Likewise, we considered the validation study to have a low risk of bias for the domains 'participants' and 'predictors' (GIMEMA V - González Rodríguez 2009 (Cabueñes coh.)), while we rated the domains 'outcome' and 'analysis' as high risk of bias due to lack of information on performance measures (Figure 18).

We rated the concern for applicability of the development study (GIMEMA D - Molica 2005 (GIMEMA cohort)) and validation study (GIMEMA V - González Rodríguez 2009 (Cabueñes coh.)) as low across all domains (Figure 15).

MDACC 2011 (Wierda 2011) (MDACC 2011 D - Wierda 2011 (MDACC))

We rated the risk of bias for the development study as high for the domain 'participants' due to inappropriate inclusion criteria (MDACC 2011 D - Wierda 2011 (MDACC)), for the 'outcome' because outcome definition did not exclude one of the predictors and for 'analysis' because of univariable selection of predictors, unclear modelling procedure and lack of performance measures. We only gave a low rating to the domain 'predictors'. Similarly, we considered the risk of bias of the validation study to be high for the domains 'outcome' for the same reason and 'analysis' due to unclear handling of missing data and low number of events (MDACC 2011 V - Molica 2016 (O-CLL1-GISL)), while we rated the domains 'predictors' and 'participants' as low risk (Figure 18).

We rated the concern for applicability of the development study as unclear for the domain 'participants' due to inappropriate inclusion criteria and low for the domains 'predictors' and 'outcome' (MDACC 2011 D - Wierda 2011 (MDACC)). We rated concern for applicability of the validation study as low across all domains (MDACC 2011 V - Molica 2016 (O-CLL1-GISL)) (Figure 15).

Morabito model (Morabito 2009) (Morabito D - Morabito 2009 (Italian cohort))

We rated the risk of bias of the development study as unclear for the domain 'participants' because it was not clear if individuals with missing data were considered (Morabito D - Morabito 2009 (Italian cohort)), low for the domains 'predictors' and 'outcome', and high for the domain 'analysis' due to low number of events, dichotomisation and univariable selection of factors and missing performance measures. In the case of the validation study (Morabito V - Gentile 2014 (O-CLL1-GISL)), we rated the risk of bias as low across all domains except 'analysis', which we considered to be of unclear risk due to lack of reporting performance measures (Figure 18).

We rated the concern for applicability of the development study as unclear for the domain 'participants' due to unclear eligibility criteria, high for the domain 'predictors' and low for the domain 'outcome' (Morabito D - Morabito 2009 (Italian cohort)). We rated as low, concern for applicability of the validation study across all domains (Morabito V - Gentile 2014 (O-CLL1-GISL)) (Figure 15).

O-CLL1 model (Gentile 2016) (O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL))

We rated the risk of bias of the development study as low across the domains 'participants' and 'predictors' (O-CLL-1D-Gentile 2016 (O-CLL-1-GISL)). We rated the domain 'outcome' as high risk of bias because one predictor was also included in the outcome definition. We rated the domain 'analysis' as high risk of bias due to unclear handling of missing data, categorisation and univariable selection of factors.

Concerning the domain 'participants', we rated one validation study as unclear due to missing eligibility criteria (O-CLL-1 V - Rani 2018 (Indian cohort)), and one as high risk due to inappropriate inclusion criteria (O-CLL1 V - Gentile 2016 (Mayo cohort)). We rated both validation studies as low risk of bias for the domain 'predictors' and high risk for the domain 'outcome' because one predictor was also included in the outcome definition. We considered the risk of bias for the domain 'analysis' to be high for one validation study due to the low number of events and no report for handling of missing data (O-CLL-1 V - Rani 2018 (Indian cohort)), while we rated the other study as unclear due to unclear handling of missing data (O-CLL-1 V - Gentile 2016 (Mayo cohort)) (Figure 18).

We rated the concern for applicability of the development study as low across all domains (O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL)). We rated concern for applicability of the two validation studies (O-CLL1 V - Gentile 2016 (Mayo cohort); O-CLL-1 V - Rani 2018 (Indian cohort)) as unclear for the domains 'participants' due to undefined eligibility criteria, the 'predictors' as low for the first and unclear for the second due to undefined timing of prognostic factor measurement, and low for both studies for the domain 'outcome' (Figure 15).

Stephens model (Stephens 2015) (Stephens TFS D - Stephens 2015 (Ohio cohort))

For risk of bias of the model for TFS in Stephens and colleagues (Stephens TFS D - Stephens 2015 (Ohio cohort)), please see description in the section for OS (above), as the authors used the same development and validation cohorts for both outcomes.

Deficiencies of prognostic model development studies (models with PROBAST rating)

Among the 12 prognostic model development studies with at least one external validation study (including the three meta-analysed models), we rated nine at high or unclear risk of bias for the domain



'analysis'. We identified the following most common issues in these studies regarding model development.

- Missing data handling: only one model development study imputed missing values; all other studies based their analysis on complete-case analyses.
- Predictor selection based on univariable analysis: eight studies derived their multivariable model on prior univariable selection using P values as a criterion.
- Correction of estimates: several development studies used splitsample or bootstrapping techniques. However, none of the studies reported a correction of their regression coefficients to reduce overfitting to the development cohort.
- Categorisation of factors: at some stage during the development process, all studies reported categorisation or dichotomisation of continuous prognostic factors.
- Model weights: 10 development studies presented a simplification of the original model formula in the form of a point score (weighting, grouping, or counting of disadvantageous prognostic factors), which results in loss of information.

Findings

Reporting of calibration was rare in the studies that we identified. If reported or provided afterwards via email, the format varied so that we were unable to summarise the 'traditional' calibration measures (calibration plots, calibration tables, O:E ratios). Therefore, we

decided to use a different way of presenting calibration graphically (Figure 19; Figure 20; Figure 21), based on something that most studies reported (survival frequencies per group). The black lines represent the development study and each coloured line one external validation cohort. The survival per risk group of the development cohort can be interpreted as the expected survival frequencies, and the survival frequencies in the external validation cohorts the observed survival frequencies. The pooled result gives the pooled observed survival frequencies.

Models with more than three external validation studies per outcome and meta-analysis

Prognostic models for the prediction of OS (with meta-analysis, > 3 external validations)

CLL-IPI (CLL-IPI D - Bahlo 2016 (development cohort))

The CLL-IPI is a prognostic score derived from univariable selection of predictors which were then entered into a forward-stepwise proportional-hazards Cox regression model with a hierarchy based on completeness of the predictor information (CLL-IPI D - Bahlo 2016 (development cohort)). The dataset was split into a training and external validation dataset after univariable selection of factors (66% and 33% of the data, respectively). Continuous factors were dichotomised based on published thresholds and quartiles. The authors integrated prognostic factors that were independently associated with the outcome in the final multivariable model in a weighted manner to construct their prognostic index (Table 2).

Table 2. Scoring of the CLL-IPI (Bahlo 2016)

Prognostic factor	Point distribution
Age (≥ 65 years)	1
Clinical stage (Rai I-IV or Binet B-C)	1
IgHV mutational status (unmutated)	2
B2-microglobulin (> 3.5 mg/L)	2
TP53 status (deleted or mutated)	4
Notes: IgHV: immunoglobulin heavy chain variable region genes	

Calibration

None of the included studies regarding the CLL-IPI reported a measure of calibration in their publication.

We included eight external validation studies of this model, with a total of 4891 individuals to calculate the pooled survival frequencies per risk group at five years (Figure 19). The pooled observed survival frequencies of all validation studies were 92.5% (89.2% to 94.8%) for the low-risk group (score 0-1), 85.0% (79.7% to 89.1%) for the intermediate-risk group (score 2-3), 64.9% (56.4% to 72.6%) for the high-risk group (score 4-6) and 40.4% (29.3%)

to 52.6%) for the very high-risk group (score 7-10). For the low, intermediate and high-risk group, the pooled result of the external validation studies approximated the survival frequencies of the model development study. In the very high-risk group, survival according to the development study would have been lower than overall in the external validation studies. For the low and intermediate-risk groups, the 95% PIs were relatively small. The other risk groups showed wide 95% PIs, indicating uncertainty (Table 3), probably due to the low number of individuals in these subgroups (N = 765 and N = 201, respectively). We judged heterogeneity (visual inspection) between external validation cohorts to be low for the calibration of the CLL-IPI.



Table 3. Survival per CLL-IPI risk group

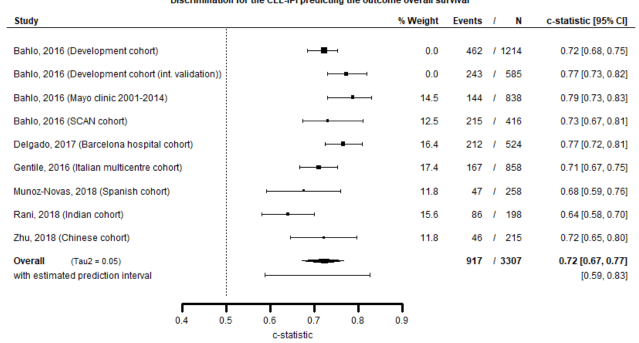
Risk group	Risk score	Development study (training dataset): percentage of persons surviving at 5 years (95% CI)	Pooled percentage of per- sons surviving at 5 years (95% CI)	95% prediction interval (PI)
Low risk	0-1	93.2% (90.5% to 96.0%)	92.5% (89.2% to 94.8%)	82.5% to 97.0%
Intermediate risk	2-3	79.3% (75.5% to 83.2%)	85.0% (79.7% to 89.1%)	68.8% to 93.6%
High risk	4-6	63.3% (57.9% to 68.8%)	64.9% (56.4% to 72.6%)	41.9% to 82.6%
Very high risk	7-10	23.3% (12.5% to 34.1%)	40.4% (29.3% to 52.6%)	13.2% to 72.4%

Discrimination

We included seven external validation studies with a total of 3307 individuals and 917 deaths observed during the overall observation time. Two studies that presented a measure for the c-statistic did not report a measure of uncertainty for this measure, which we estimated using the Newcombe method (CLL-IPI V - Gentile 2016)

(Italian cohort); CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort)). For one study (CLL-IPI V - Zhu 2018 (Chinese cohort)), we included the AUC as a measure of discrimination instead of the c-statistic. The pooled c-statistic of the seven studies was 0.72 (95% CI 0.67 to 0.77), with some degree of heterogeneity (between-study standard deviation, tau = 0.21). The 95% PI ranged from 0.59 to 0.83 (Figure 22).

Figure 22. Meta-analysis of the c-statistic for the CLL-IPI model (Bahlo 2016)



Discrimination for the CLL-IPI predicting the outcome overall survival

The c-statistic of the model development study was 0.72 (95% CI 0.68 to 0.75). Based on sensitivity analysis omitting the study that used the AUC instead of the c-statistic (Figure 23), and comparing estimated and reported 95% CI of the c-statistic (Figure 24; Figure

25), we conclude that this had no effect on the result. Four out of seven external validation studies replaced the prognostic factor TP53 with a proxy, del(17p); sensitivity analysis showed no substantial differences in discriminative performance (Figure 26).



Figure 23. Sensitivity analysis of the c-statistic for the CLL-IPI excluding the study reporting AUC

Sensitivity analysis AUC: Discrimination for the CLL-IPI predicting the outcome overall survival

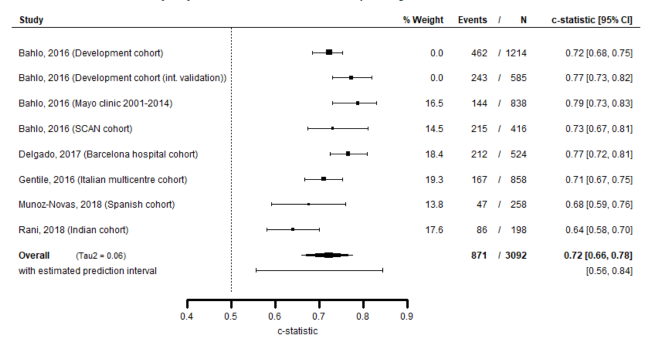


Figure 24. Sensitivity analysis of the c-statistic for the CLL-IPI excluding the study reporting no 95% CI

Sensitivity analysis complete data: Discrimination for the CLL-IPI predicting the outcome overall survival

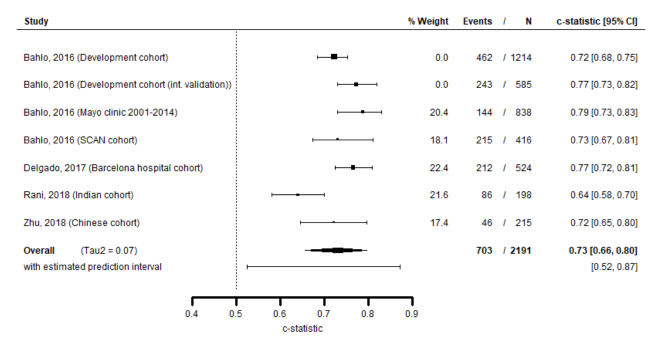




Figure 25. Sensitivity analysis of the c-statistic for the CLL-IPI excluding the study reporting no 95% CI. All 95% CIs were replaced by estimates using the Newcombe method for a comparison with Figure 27

Sensitivity analysis Newcombe: Discrimination for the CLL-IPI predicting the outcome overall survival

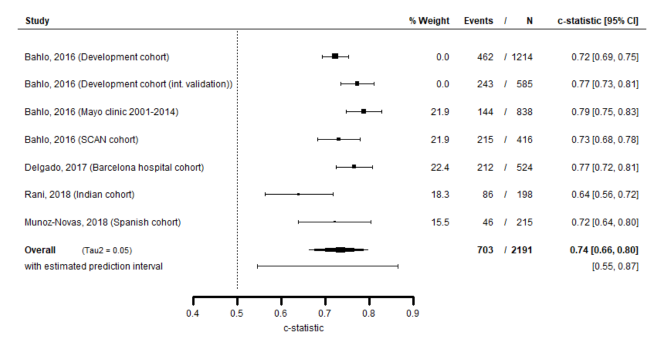
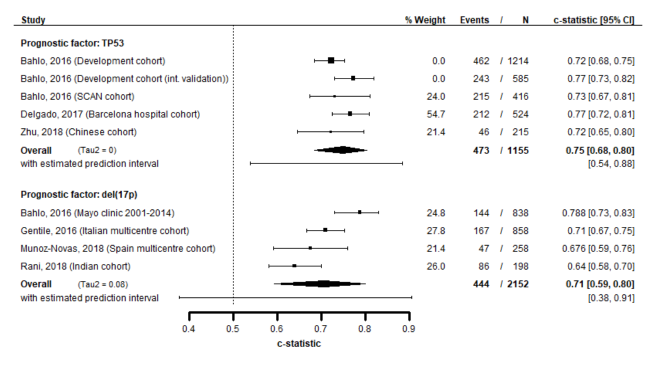


Figure 26. Sensitivity analysis of the c-statistic for the CLL-IPI regarding the availability of the predictor TP53 and its proxy del(17p)

Sensitivity analysis TP53: Discrimination for the CLL-IPI predicting the outcome overall survival





Barcelona-Brno model (Barcelona-Brno D - Delgado 2017 (Barcelona cohort))

The Barcelona-Brno score is a prognostic score constructed by comparing the best combinations of the five factors included in another model for CLL (Barcelona-Brno D - Delgado 2017 (Barcelona cohort)), the CLL-IPI by Bahlo 2016 CLL-IPI D - Bahlo 2016 (development cohort)). Incomplete data were most likely part of the exclusion criteria, and therefore persons with missing data were excluded from the analysis in the primary publication. The score classified individuals into three risk groups: persons without del(11q) or del(17p) and mutated IgHV status were classified as low risk; persons with del(11q) or del(17p) and unmutated IgHV status were classified as high risk; all other persons were classified as intermediate risk.

Calibration

None of the included studies in reference to the Barcelona-Brno model reported a measure of calibration in their publication.

We included three external validation studies of this score with a total of 1974 individuals to calculate the pooled survival frequencies per risk group at five years (Figure 20). The pooled observed survival frequencies of all validation studies were 90.5% (95% CI 85.1% to 94.0%) for the low-risk group, 79.7% (95% CI 70.7% to 86.5%) for the intermediate-risk group and 62.5% (95% CI 49.3% to 74.1%) for the high-risk group. For the low and intermediaterisk group, the pooled result of the external validation studies approximated the survival frequencies of the model development study. In the high-risk group, survival according to the development study would have been higher than overall in the external validation studies. For the low-risk group, the 95% PI was relatively small. However, the intermediate and high-risk groups showed a wide 95% PI, indicating uncertainty (Table 4). We judged heterogeneity (visual inspection) between external validation cohorts to be low for the calibration of the Barcelona-Brno score.

Table 4. Survival per Barcelona-Brno model risk group

Risk group	Risk group	Development study: percentage of persons surviving at 5 years (95% CI)	Pooled percentage of persons surviving at 5 years (95% CI)	95% prediction interval (PI)
Low risk	No del(11q) or del(17p) and mutated IgHV status	93.4% (90.4% to 96.5%)	90.5% (85.1% to 94.0%)	80.4% to 95.7%
Intermediate risk	All other	83.8% (78.4% to 89.6%)	79.7% (70.7% to 86.5%)	63.1% to 90.0%
High risk	Del(11q) or del(17p) and unmutated IgHV status	70.0% (59.3% to 82.7%)	62.5% (49.3% to 74.1%)	41.3% to 79.7%

Notes: IgHV: immunoglobulin heavy chain variable region genes

$\underline{\text{Discrimination}}$

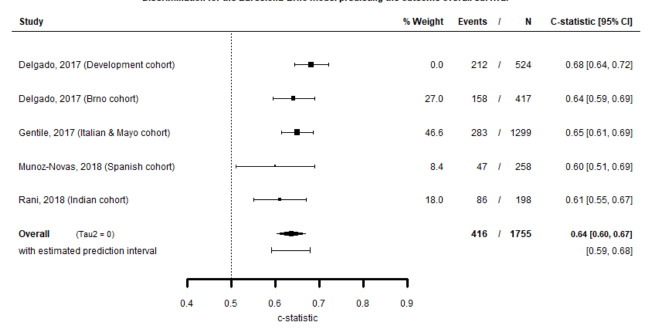
For our meta-analysis of the c-statistic, we included four external validation studies with a total of 1755 individuals and 416 deaths observed during the overall observation time. Two studies that presented a measure for the c-statistic did not report a measure

of uncertainty, which we estimated using the Newcombe method (Barcelona-Brno V - Gentile 2017 (Italian & Mayo); Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.)). The pooled c-statistic of the four studies was 0.64 (95% CI 0.60 to 0.67), with no heterogeneity between studies (tau = 0.00). The 95% PI ranged from 0.59 to 0.68 (Figure 27).



Figure 27. Meta-analysis of the c-statistic for the Barcelona-Brno model (Delgado 2017)

Discrimination for the Barcelona-Brno model predicting the outcome overall survival



The c-statistic of the model development study was 0.68 (95% CI 0.64 to 0.72); it was unclear whether bootstrapping techniques were used to obtain this estimate.

MDACC 2007 (Wierda 2007) (MDACC 2007 D - Wierda 2007 (MDACC))

The MDACC 2007 model is a prognostic model based on univariable selection of predictors that were, when significant, entered into

a proportional-hazards Cox regression model (MDACC 2007 D - Wierda 2007 (MDACC)). Bootstrapping was used to account for optimism in the c-statistic. The authors provided the precise formula to calculate a point score which could be translated graphically by a nomogram into individual survival probabilities at five or 10 years. In addition, for simplified use in clinical practice, the authors created an index score in a weighted manner (Table 5).

Table 5. Scoring of the MDACC 2007 model

	Point distribution			
Prognostic factor	0	1	2	3
Age (years)		< 50	50-65	> 65
ß-2 microglobulin (mg/L)	< ULN	1-2 × ULN	> 2 × ULN	
ALC (× 10 ⁹ /L)	< 20	20-50	> 50	
Gender	Female	Male		
Rai stage	0-11	III-IV		
No. of involved nodal groups	≤2	3		
Notes: ULN: upper limit of normal; ALC: acute ly	mphocyte count			



Calibration

The development study provided a calibration plot based on bootstrap sampling. The data points representing the observed versus expected survival probabilities are close to the ideal line. At the higher frequencies, the nomogram slightly underestimates survival. Measures of calibration were not reported in any of the external validation studies.

We included five external validation studies of this model with a total of 3786 individuals for calculating the pooled survival frequencies per risk group at 5 years (graphically represented in Figure 21). The pooled observed survival frequencies of all validation studies was 97.0% (95% CI 94.3% to 98.4%) for the low-risk group (score 1-3), 82.3% (74.6% to 88.0%) for the intermediate-risk group (score 4-7) and 45.6% (31.3% to 60.5%) for the high-risk group (score \geq 8). For the low and intermediate-risk group, the pooled result of the external validation studies approximated the survival frequencies of the model development study. In the high-risk group, survival according to the development study would have been higher than overall in the external validation studies. For the low-risk group, the 95% PI was relatively small. The intermediate and high-risk group showed wide 95% PIs, indicating uncertainty (Table 6). We judged heterogeneity (visual inspection) of calibration between external validation cohorts to be low for the MDACC 2007 index score.

Table 6. Survival per MDACC 2007 model risk group

Risk group	Index score	Development study: Percentage of persons surviving at 5 years (95% CI)	Pooled percentage of persons surviving at 5 years (95% CI)	95% prediction interval (PI)
Low risk	1-3	97.0% (95.0% to 99.0%)	97.0% (94.3% to 98.4%)	90.9% to 99.0%
Intermediate risk	4-7	80.0% (78.0% to 82.0%)	82.3% (74.6% to 88.0%)	61.5% to 93.1%
High risk	≥ 8	55.0% (47.2% to 62.8%)	45.6% (31.3% to 60.5%)	21.2% to 72.3%

Discrimination

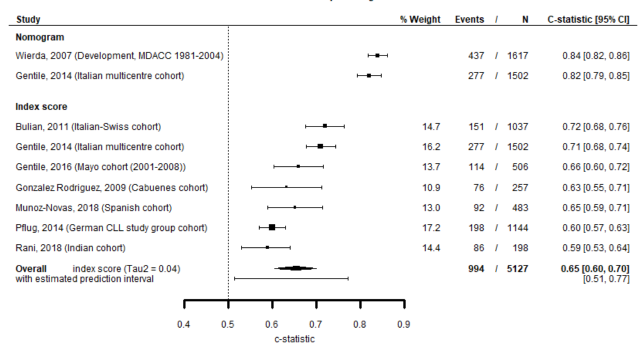
We included seven external validation studies with a total of 5127 individuals and 994 deaths observed during follow-up for meta-analysis of the c-statistic in the case of the index score. Only one external validation study provided sufficient data on performance for the nomogram, which is presented together with the development study discrimination on the upper part of the forest plot (Figure 28). Four studies that reported a measure for the c-statistic did not report a measure of uncertainty, which we estimated using the Newcombe method (MDACC 2007 V - Bulian

2011 (Italian-Swiss); MDACC 2007 V - Gentile 2014 (Italian cohort); MDACC 2007 V - Gentile 2016 (Mayo cohort); MDACC 2007 V - Muñoz-Novas 2018 (Spanish cohort))). For one study (MDACC 2007 V - González Rodríguez (Cabueñes), we included the AUC as measure of discrimination instead of the c-statistic. The pooled estimate of the seven studies was 0.65 (95% CI 0.60 to 0.70), with some degree of heterogeneity (between-study standard deviation, tau = 0.21). The prediction interval, which describes a range for the predicted model discrimination in a new validation study of the model, ranged from 0.51 to 0.77.



Figure 28. Meta-analysis of the c-statistic for the MDACC 2007 model (Wierda 2007)

Discrimination for the MDACC 2007 model predicting the outcome overall survival



The bootstrap-corrected c-statistic of the model development study was 0.84 (95% CI 0.82 to 0.86). The validation study for the nomogram showed a similar discrimination of 0.82 (95% CI 0.79 to 0.85). Other studies that used the nomogram provided the point score of the formula. However, due to the poor graphical quality of the presented nomogram, authors did not attempt to estimate the expected individual survival probabilities per individual (MDACC 2007 V - Bulian 2011 (Italian-Swiss); MDACC 2007 V - Gentile 2014 (Italian cohort)).

Prognostic models for the prediction of PFS or TFS (with metaanalysis, > 3 external validations)

We did not identify any prognostic model development studies that were externally validated more than three times for the outcomes PFS, TFS or TFS.

Models with one to three external validation studies

Prognostic models for the prediction of OS (without metaanalysis, 1 to 3 external validations)

GCLLSG model (Pflug 2014) (GCLLSG D - Pflug 2014 (GCLLSG))

The GCLLSG model was derived from univariable selection of predictors which were entered into a forward and backward stepwise proportional-hazards Cox regression model (GCLLSG D - Pflug 2014 (GCLLSG)). The authors controlled for several factors specific to the RCT data that were used for model-building (e.g. study, treatment etc). Bootstrapping techniques were used to test the robustness of the Cox regression model. Persons with missing data were excluded from analysis. Factors were categorised based on published thresholds and quartiles. The authors assigned risk scores to factors, which proved to be independent in the final multivariable model, in a weighted manner to construct the prognostic index presented in Table 7.

Prognostic factor	Point distribution
FISH category del(17p)	6
Serum thymidine kinase (> 10.0 U/L)	2
B2-microglobulin (> 3.5 mg/L)	2
B2-microglobulin (> 1.7 and ≤ 3.5 mg/L)	1



IgHV mutational status (unmutated)	1
ECOG PS (> 0)	1
FISH category del(11q)	1
Gender (male)	1
Age (> 60 years)	1

Notes: FISH: fluorescence in situ hybridisation; IgHV = immunoglobulin heavy chain variable region genes; ECOG PS: Eastern Cooperative Oncology Group Performance Status

Four risk groups were determined: low risk (0-2 points), intermediate risk (3-5 points), high risk (6-10 points) and very high risk (> 10 points). Five- and six-year OS was reported for each point score.

Calibration

Calibration was not reported for the development and external validation study (GCLLSG V - Pflug 2014 (Mayo cohort)).

In the development study, survival was 95.2%, 86.9%, 67.6% and 18.7% at five years for the low, intermediate, high and very highrisk group, respectively. In the validation study, survival was 95.2%, 91.1%, 71.7% and 13.6% at five years for the low, intermediate, high and very high-risk group, respectively.

Discrimination

The c-statistic of the prognostic score in the model development study was 0.75 (95% CI 0.70 to 0.78), discrimination in the external validation cohort was 0.77 (95% CI 0.70 to 0.83).

Rossi (Rossi 2013) (Rossi D - Rossi 2013 (Italian cohort))

The prognostic score by Rossi and colleagues was developed in several steps (Rossi D - Rossi 2013 (Italian cohort)). Firstly, the authors identified factors that were independently associated with OS by univariable and multivariable Cox regression analysis. Bootstrapping techniques were used to test the robustness of the model. In the next step, the factors were entered into a decision tree algorithm to divide individuals in subgroups. To test the stability of the decision tree, the random survival forest method with amalgamation algorithm was used. It was unclear how missing data were handled.

Calibration

No study reported a measure of calibration for this model.

In the development study, OS at five years was 86.9% in the very low-risk group (del13q14 only), 77.6% in the low-risk group (normal/+12), 65.9% in the intermediate-risk group (NOTCH1 and/ or SF3B1 mutations and/or del11q22-q23 in the absence of TP53 and BIRC3 abnormalities) and 50.9% in the high-risk group (TP53 disruption and/or BIRC3 disruption independent of co-occurring lesions). One external validation study reported median survival per risk group (not reached for the very low-risk group, 13.8 years for the low-risk group, 11.2 years for the intermediate-risk

group and 7.7 years for the high-risk group) (Rossi V - Rossi 2013 (unclear)). In the other validation study, OS at five years was 91% in the very low-risk group, 90% in the low-risk group, 75.2% in the intermediate-risk group and 62.1% in the high-risk group (Rossi V - Jeromin 2014 (Munich cohort)).

Discrimination

The c-statistic of the prognostic score in the development cohort was 0.64. Only one of the two validation studies reported a measure of discrimination; the c-statistic was 0.66 (95% CI not reported).

Stephens model (Stephens 2015) (Stephens OS D - Stephens 2015 (Ohio cohort))

To develop prognostic models for OS, the authors identified significant prognostic factors in multivariable proportional hazards models using backwards selection. Based on the final model, they defined a simplified risk score based on the strength of the association. Missing data were accounted for by multiple imputation techniques. The model and score for OS included the factors ECOG performance status, age and lactate dehydrogenase (LDH).

Calibration

No study reported a measure of calibration for the prognostic models.

In the development cohort, the percentage of alive persons at two years was 89% (95% CI 74% to 96%) for score 0, 66% (95% CI 41% to 82%) for score 2 and 0% for score 4. In the validation cohort, the percentage of alive persons at two years was 95% (95% CI 83% to 99%) for score 0, 80% (95% CI 55% to 92%) for score 2 and 20% (95% CI 1% to 58%) for score 3.

Discrimination

The discrimination of the exact model for OS was 0.76 (P < 0.03) and 0.73 (P < 0.0001) for the development cohort. The validation cohort showed a c-statistic of 0.68 for the simplified score for OS.



Prognostic models for the prediction of PFS or TFS (without meta-analysis, 1 to 3 external validations)

Baliakas model (Baliakas 2019) (Baliakas D - Baliakas 2019 (multicentre))

The model presented by Baliakas and colleagues was derived from univariable selection of predictors which were entered into a proportional-hazards Cox regression model (Baliakas D - Baliakas 2019 (multicentre)), that was internally validated by bootstrapping and further confirmed by recursive partitioning based on conditional inference trees and merging of terminal nodes by an amalgamation algorithm. First, the authors split individuals in two groups, mutated and unmutated IgHV. The risk groups for mutated cases were as follows: (1) low risk: non-TP53 abnormality/ +12/subset #2 membership and Binet stage A; (2) intermediate risk: Binet A with one of the following: TP53 abnormality and/or +12 and/or subset #2 membership, (3) high risk: Binet B, (4) very high risk: Binet C. The risk groups for unmutated cases were as follows: (1) very low risk: non-TP53abn/SF3B1mut/del(11q) female Binet A; (2) low risk: non-TP53abn/SF3B1mut/del(11q) male Binet A; (3) intermediate risk: Binet A with one of the following: TP53abn and/ or SF3B1mut and/or del(11q); (4) high risk: Binet B; (5) very high risk: Binet C.

Calibration

No measures of calibration were reported for this model.

In the development study, the five-year treatment probability for mutated cases was 12% for group 1, 40% for group 2, 64% for group 3 and 92% for group 4. For unmutated cases, the five-year treatment probability was 45% for group 1, 65% for group 2, 78% for group 3, 90% for group 4 and 100% for group 5.

In the external validation cohort, the five-year treatment probability was only reported for the intermediate-risk groups (for mutated cases 43%, group 2; and for unmutated cases 74%, group 3).

Discrimination

In the development cohort, the discrimination was 0.745 (SE = 0.013) and 0.753 (SE = 0.013) for the cases with mutated IgHV and unmutated IgHV status, respectively. Discrimination was not reported for the external validation cohort.

GIMEMA model (Molica 2005) (GIMEMA D - Molica 2005 (GIMEMA cohort))

The GIMEMA model was derived from univariable selection of predictors which were entered into a proportional-hazards Cox regression model (GIMEMA D - Molica 2005 (GIMEMA cohort)). No internal validation was reported. Negative risk factors that were independent in multivariable analysis (lymphocyte doubling time < 12 months, lymphocyte count >30 \times 10 9 per L and Rai stage I to II) were assigned one point each (not weighted for HR) to create a score. Gender was added afterwards as relevant for individuals with score 0, but not for individuals with scores 1 to 3. Three risk groups were created: (1) females with score 0; (2) males with score 0; and (3) individuals with score 1 to 3 of either gender.

Calibration

No measure of calibration was reported for this model.

In the development study, the 10-year PFS was 76.2% for group 1, 61.4% for group 2 and 37.8% for group 3, respectively. In the validation study, the 10-year treatment-free survival was 89% for group 1, 54% for group 2 and 0% for group 3 (GIMEMA V - González Rodríguez 2009 (Cabueñes coh.)).

Discrimination

The development study did not report a measure of discrimination. The AUC for the validation study was 0.58 (95% CI 0.49 to 0.66).

MDACC 2011 (Wierda 2011) (MDACC 2011 D - Wierda 2011 (MDACC))

The MDACC 2011 model (MDACC 2011 D - Wierda 2011 (MDACC)) is a prognostic model derived from univariable selection of predictors. After univariable selection, the dataset was split into a training and test set. Significant predictors entered into a proportional-hazards Cox regression model (forward selection procedure with P < 0.1, and removed if not P < 0.5 in final model) were developed on the test set and applied to the test set. The factors of the final model were entered into Cox regression models for 1000 bootstrap samples to show their robustness. The final model formula can be used to derive a point score that can be translated graphically by a nomogram into individual survival probabilities at five or 10 years.

The formula is as follows: [I(No. of lymph node sites involved = 3) \times 7.370 + I(FISH11q del) \times 9.312 + I(FISH 17p del) \times 11.285 + (diameter of largest cervical lymph node in cm) \times 4.172 + (LDH/100) \times I([IgHV gene mutated] \times 5.000 + (LDH/100) \times I(IgHV gene = unmutated) \times 1.065] + 35.467.

Calibration

No study reported a measure of calibration for this model.

Discrimination

The development study did not report a measure of discrimination. The validation study included a total of 337 participants (91 treated) and reported a c-statistic of 0.71 (95% CI 0.60 to 0.82) (MDACC 2011 V - Molica 2016 (O-CLL1-GISL)).

Morabito model (Morabito 2009) (Morabito D - Morabito 2009 (Italian cohort))

The prognostic model developed by Morabito and colleagues (Morabito D - Morabito 2009 (Italian cohort)) was derived from univariable analysis, including dichotomisation of factors with cutoff determination by ROC curves. Significant (P > 0.05) prognostic factors were entered into proportional-hazards Cox regression models. Each unfavourable marker that remained significant was assigned one point, and the sum of these points formed the final score (CD38 positive = 1, ZAP-70 positive = 1, IgHV unmutated status = 1). Individuals with scores of two or three points were categorised together to form the high-risk group.

Calibration

No study reported a measure of calibration for this model.

The five-year PFS of the validation study for this model was 91.7% for the low-risk group (score 0), 82.9% for the intermediate-risk group (score 1) and 57.4% for the high-risk group (score 2-3) (Morabito V - Gentile 2014 (O-CLL1-GISL)).

Discrimination



Both the development and validation study did not report a measure of discrimination.

O-CLL1 model (Gentile 2016) (O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL))

The prognostic score developed by Gentile and colleagues was derived from univariable selection of predictors (O-CLL-1 D-Gentile 2016 (O-CLL-1-GISL)). Before entering the factors into proportional-hazards Cox regression models (P < 0.05), continuous factors

were dichotomised using established thresholds or ROC curves. Bootstrapping techniques were used to test the robustness of the Cox regression model. There were no persons with missing values in the sample. The authors assigned risk scores to each prognostic factor that was significant in multivariable analysis based on their hazard ratio (Table 8). Individuals were divided into three different risk categories for TFS, low (score 0 to 2), intermediate (score 3 to 5) and high risk (score 6 to 7).

Table 8. Scoring of the O-CLL1 model

	Point distribution		
Prognostic factor	0	1	2
Rai staging system	0	1-11	-
B2-microglobulin	Normal	-	Elevated
ALC (10 ⁹ /L)	< 10	-	≥ 10
IgHV mutational status	Mutated	-	Unmutated

Notes: ALC: acute lymphocyte count; IgHV: immunoglobulin heavy chain variable region genes

Calibration

No study reported a measure of calibration for this model.

In the development study, treatment-free survival at three years was 95.3% in the low risk, 74.5% in the intermediate-risk and 28.6% in the high-risk group. Outcome frequencies were also provided per risk score. One external validation study showed similar percentages of treatment-free survival at three years with 95.5%, 78.9% and 40.6% for the low, intermediate and high-risk group, respectively (O-CLL1 V - Gentile 2016 (Mayo cohort)). The other external validation study presented results in the form of a Kaplan-Meier curve (O-CLL-1 V - Rani 2018 (Indian cohort)).

Discrimination

The c-statistic of the prognostic score in the model development study was 0.75 (P < 0.001). The c-statistic in the external validation cohorts were 0.72 (P < 0.001) (O-CLL1 V - Gentile 2016 (Mayo cohort)), and 0.55 (95% CI 0.49 to 0.60) (O-CLL-1 V - Rani 2018 (Indian cohort)).

Stephens model (Stephens 2015) (Stephens TFS D - Stephens 2015 (Ohio cohort))

To develop prognostic models for TFS, the authors identified significant prognostic factors in multivariable proportional hazards models using backwards selection. Based on the final model, they defined a simplified risk score based on the strength of the association. Missing data were accounted for by multiple imputation techniques. The model and score for TFS include the factors ECOG performance status, Rai stage, age, white blood cell count (WBC) and FISH category del(11q22.3). Risk groups were summarised as 0 to 1 points, 2 to 3 points and ≥ 4 points.

Calibration

No study reported a measure of calibration for the prognostic model for TFS.

In the development cohort, the percentage of persons who were treatment-free at two years was 85% (95% CI 0.60 to 0.95) for the low-risk group, 51% (95% CI 32 to 67) for the intermediate-risk group, and 0% for the high-risk group. In the validation cohort, the percentage of persons who were treatment-free at three years was 63% (95% CI 39 to 79) for the low-risk group, 26% (95% CI 15 to 39) for the intermediate-risk group and 16% (95% CI 6 to 29) for the high-risk group.

Discrimination

The discrimination of the exact model for TFS was 0.84 (P < 0.017) for the development cohort. The validation cohort showed a c-statistic of 0.66 for the simplified score for TFS.

Models without any external validation studies

The main characteristics of prognostic models for which no external validation studies could be found are described in Appendix 7.

Subgroup differences

We did not explore subgroup differences as the number of external validation studies per model with available data were too small. Three of the four external validation studies for the Barcelona-Brno model defined CLL by the International Workshop on CLL Guideline (Hallek 2008), and one did not report diagnostic criteria. Among the seven external validations of the CLL-IPI, three studies used the International Workshop on CLL Guideline (Hallek 2008), and four did not clearly refer to the diagnostic criteria or used a combination



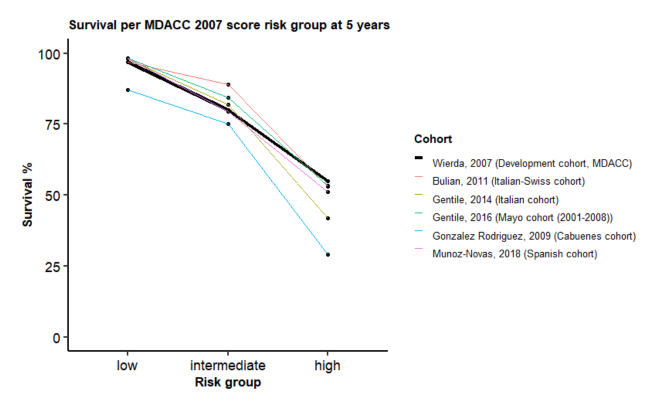
of criteria. One external validation study of the MDACC 2007 model used the International Workshop on CLL Guideline (Hallek 2008), two studies' cohorts were based on the NCI working group criteria (Cheson 1996), and four studies did not clearly report the criteria or used a combination of criteria.

Sensitivity analysis

We planned to explore the effect of risk of bias on the performance measures for the external validation studies per model. We were not able to conduct this analysis as we rated nearly all studies as high or unclear risk of bias, leaving no studies with a low risk of bias for inclusion in the sensitivity analysis.

One external validation study for the CLL-IPI reported the AUC instead of the c-statistic (CLL-IPI V - Zhu 2018 (Chinese cohort)), which we included in our meta-analysis. The pooled performance estimate for discrimination did not change (0.72, 95% CI 0.67 to 0.77 including the study; 0.72, 95% CI 0.66 to 0.78 excluding the study; see Figure 23). For the MDACC 2007 model, we also included one AUC instead of the c-statistic in the analysis. By removing this study (MDACC 2007 V - González Rodríguez (Cabueñes)), the pooled performance estimate for discrimination barely changed (from 0.65 to 0.66, 95% CI 0.60 to 0.71; for illustration, see Figure 29).

Figure 29. Sensitivity analysis of the c-statistic for the MDACC 2007 model excluding the study reporting AUC



To examine the effect of estimating the 95% CIs according to the Newcombe method, we conducted a sensitivity analysis exemplary for one included prognostic model, the CLL-IPI. In Figure 24, we limited the meta-analysis of the c-statistic to all studies that reported the 95% CI, which did not change the overall pooled estimate and 95% CIs substantially (from 0.72, 95% CI 0.67 to 0.77 for the main analysis to 0.73, 95% CI 0.66 to 0.80 for the analysis limited to studies with 95% CI reported). We calculated the 95% CIs based on Newcombe for all studies and the pooled results to compare this extreme scenario with the scenario that all 95% CIs were provided. Figure 25 shows that the estimation did not change the 95% CIs. The 95% PI changed negligibly.

For the CLL-IPI, one of the predictors was commonly missing (TP53 mutation) and replaced by a proxy prognostic factor (del(17p)). To see if the replacement limited discriminative performance, we conducted separate meta-analyses (Figure 26). The 95% CI of the pooled c-statistic overlapped, however, the number of studies in each analysis was very small (N = 3 and N = 4).

DISCUSSION

Summary of main results

We aimed to identify and describe all prognostic models for untreated CLL and their corresponding external validation studies. We identified 12 prognostic models with at least one external validation (Appendix 6). Of these, three models were validated externally more than three times, with appropriate reporting to allow us to conduct a meta-analysis of the c-statistic, which is a measure of discriminative performance of a model. Additionally, we identified 40 models or scores without any external validation study, which are further described in Appendix 7 and Characteristics of studies awaiting classification.



Results for models with more than three external validation studies per outcome for meta-analysis

Prognostic models for the prediction of OS (with meta-analysis, > 3 external validations)

The pooled c-statistic of the CLL-IPI, a point score calculated from five factors, was based on seven external validation studies with a total of 3307 individuals. The pooled estimate of its discriminative performance in seven external validation studies was 0.72 (95% CI 0.67 to 0.77). The 95% PI ranged from 0.59 to 0.83, indicating that from 100 external validation studies for this model, 95 will show a c-statistic within this range. Calibration was not reported in the publications, and thus the prognostic score was not updated based on calibration. Studies showed somewhat heterogeneous results regarding the pooled survival frequencies (observed survival frequencies) as compared to the development study survival frequencies (expected survival frequencies). Due to our diverging representation of calibration, to make assumptions regarding the calibration of the models, we assumed that the survival frequency of the development cohort can be interpreted as our expected frequency. The pooled survival per risk group was higher than observed in the development study for the very highrisk group, indicating that the model underpredicted survival in this group. This may be explained by the fact that the model was developed using data including a higher proportion of individuals with treatment indication at recruitment, who are participants with a worse prognosis than the general population with CLL. The curves showed a relatively homogeneous pattern regarding the differences between risk groups, although frequencies varied more widely for the very high-risk group, probably due to the low number of participants.

The pooled c-statistic of the Barcelona-Brno score, which contains two of the prognostic factors also included in the CLL-IPI, was derived from four external cohorts with a total of 1974 individuals. The pooled c-statistic was 0.64 (95% CI 0.60 to 0.67) and the 95% PI ranged from 0.59 and 0.68. Results between cohorts were somewhat heterogeneous, although the number of studies was relatively low to draw meaningful conclusions. Calibration was not reported in the publications, and thus the prognostic score was not updated based on calibration. The pooled survival per risk group was lower than observed in the development study, especially for the high-risk group, which means that across cohorts, the model overestimated survival as compared to the development cohort. This judgement is mainly driven by the Czech cohort. The pattern was consistent across studies.

The pooled c-statistic of the MDACC 2007 index score, which includes six prognostic factors, was derived from seven external validation studies with a total of 5127 individuals. The pooled c-statistic was 0.65 (95% CI 0.60 to 0.70; 95% PI: 0.51 to 0.77); discrimination was somewhat heterogeneous between studies. Calibration was not reported in any of the studies validating the index score, thus, no model updates were reported. The pooled survival per risk group matched the survival in the development study for the low and intermediate-risk groups. In the high-risk group, the pooled survival per risk group was lower than observed in the development study, thus the model overestimated the survival of the high-risk group. The pattern was consistent across studies.

From these three prognostic models for untreated individuals with CLL included in meta-analysis, the CLL-IPI performed best regarding the discrimination between persons with a 'as good as' compared to a worse prognosis. The score underpredicted survival for the very high-risk group, which was a small group of individuals defined by TP53 mutation or deletion.

The three models consist of unweighted (Barcelona-Brno D - Delgado 2017 (Barcelona cohort)) or weighted (CLL-IPI D - Bahlo 2016 (development cohort), MDACC 2007 D - Wierda 2007 (MDACC)) simplified point scores, which implies that information concerning the correct weighting of the prognostic factors was lost during simplification. Instead of individual outcome frequencies, the models result in a score on a limited scale, which reduced the possible outcome range compared to the original formula. Consequently, the simplification may have resulted in a reduction of the discriminative performance.

Prognostic models for the prediction of PFS or TFS (with metaanalysis, > 3 external validations)

We did not identify any prognostic model development studies that were externally validated more than three times for the outcomes PFS, TFS or TFS.

Models with one to three external validation studies

Prognostic models for the prediction of OS (without metaanalysis, 1 to 3 external validations)

In total, three models were developed to predict OS (GCLLSG D - Pflug 2014 (GCLLSG); Rossi D - Rossi 2013 (Italian cohort); Stephens OS D - Stephens 2015 (Ohio cohort)), with one, two and one external validation study, respectively. Due to the limited information on the predictive performance outside the model development setting, we would not yet consider these models ready for use in clinical practice.

Prognostic models for the prediction of PFS or TFS (without meta-analysis, 1 to 3 external validations)

In total, one model was developed to predict PFS (GIMEMA D - Molica 2005 (GIMEMA cohort)) and five were developed to predict TFS (Baliakas D - Baliakas 2019 (multicentre); MDACC 2011 D - Wierda 2011 (MDACC); Morabito D - Morabito 2009 (Italian cohort); Stephens TFS D - Stephens 2015 (Ohio cohort); O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL)), with one external validation study per model for the first five and two for the last mentioned. Due to the limited information on the predictive performance outside the model development setting, we would not yet consider these models ready for use in clinical practice.

Models without any external validation studies

An abundance of prognostic models and scores without any application in external cohorts has been published, with a varying degree of reporting quality. Without any further knowledge on their performance in different settings than the development cohort, we would not recommend their use for patients.



Overall completeness, certainty of the evidence and study limitations of externally validated models

Overall completeness of the data

As expected from results of previous research (Heus 2018), reporting of the included models and their validation studies was poor. We have summarised the missing information per model for the development study and external validation studies for the primary model outcome in Appendix 8.

For the CLL-IPI, none of the studies reported calibration (although we obtained information for five studies from corresponding authors). Reporting of discrimination was higher (8 out of 11, 6 times with 95% CI).

For the Barcelona-Brno score, none of the studies reported calibration (although we obtained information for one study from the corresponding authors). Reporting of discrimination was slightly higher (5 times out of 6, 3 times with 95% CI).

For the MDACC 2007 index score, two of the studies reported calibration. Reporting of discrimination was slightly higher (8 times out of 10, 7 times with 95% CI).

Of the other nine models with 17 validation studies that were not meta-analysed, none reported calibration (we obtained information on calibration for one study from the corresponding authors). Reporting of discrimination was higher (12 times, seven times with 95% CI).

Overall, across all categories of included studies (development studies with and without external validations, external validations), information on calibration and discrimination was lacking. Calibration was reported in six studies and discrimination in 37 out of 85 studies.

Many studies either excluded persons with missing prognostic factor data at enrolment, or excluded persons with missing data from the analysis. In most cases, we do not have information on the number of persons in the complete database or sample, and a comparison between included and excluded persons is rarely provided. Instead of removing data of individuals with missing data completely, multiple imputation would be preferable, but should be interpreted carefully (Moons 2019; Steyerberg 2014).

None of the studies explicitly mentioned reasons for censoring, a common phenomenon in time-to-event data, in their cohort which may be particularly relevant for studies based on RCT data.

Certainty of the evidence

At the date of submission of this review, no official GRADE guidance for grading the summarised results of meta-analysis of prognostic models was available. Hence, we refrained from rating the certainty of the evidence.

Study limitations of prognostic model development studies

The majority of PROBAST ratings for model development studies was high. The main reason for this was handling missing data, predictor selection based on univariable selection of predictors, failure to correct estimates for optimism, categorisation of prognostic factors and simplification of models.

Potential bias in the review process

To prevent bias in the review process, we performed all relevant steps in duplicate and solved discrepancies in group discussions. We developed a sensitive search strategy and tracked the references and citations of all included references. However, we did not search conference abstracts or trial registries, since conference abstracts do not provide sufficient information for 'Risk of bias' assessment and prognostic studies are rarely prospectively registered, but rather built on existing retrospective databases. Thus, we would not expect to find any relevant studies by searching those additional databases.

As this is a new review type, our methods evolved during the process. We decided to adapt the newest methods regardless of the protocol, which may have introduced bias.

- Prespecification of prognostic factors: we did not limit our inclusion to models with e.g. a minimum set of clinically relevant factors or a specific kind of factor (such as non-invasive or genetic factors only). Therefore, we made no distinction regarding which predictors were included in the retrieved models we included all developed and validated models that were the subject of our review, regardless of which type of predictors they included. We did not aim to rate the individual factors, examine the relationship between any one factor with our outcomes or the strength of any one factor above other factors, as this would be the subject of a prognostic factor review.
- Search: we have slightly adapted the search to make it more inclusive, and extended it to include studies published since the inception of MEDLINE instead of using the cut-off year 1990. Based on the recommendation of the Cochrane fast-track service, we have added a search strategy for Embase. We believe that screening more references has not introduced bias.
- Screening: the definition of prognostic models and the reporting standards and, with that, the amount of information expected in a publication has changed substantially over time. To avoid bias, we have included all studies that explicitly stated the aim of developing a score for prognostication, although from today's point of view, we would not consider these scores a well-developed prediction model, because they did not follow the recommended steps in the model-building process. Most of these studies can be found in Appendix 7, as models without external validation studies.
- Risk of bias: since publication of the protocol, PROBAST was published. In our protocol, we did not specify the data analysis process. To avoid bias and use the most recently developed methods, we worked in close collaboration with the Cochrane Prognosis Methods group, and some authors of PROBAST are coauthors of this review. For transparency of our group decisions, we added the text under 'Solving disagreements'.
- Analysis: we planned to summarise the performance measures
 for calibration and discrimination. In the review we specified,
 according to most recent developments, that we would
 summarise the external validation studies per model. For some
 studies, we estimated the 95% CI for the c-statistic due to lack of
 reporting. Based on our sensitivity analysis, we assume that this
 estimation has not introduced bias.

We did not assess the likelihood of publication bias, i.e. a preferential publication of studies which show better model



performance, because currently, there is no established standard for this.

We would like to mention that external validation studies are ideally carried out by independent researchers (using independent samples). For the studies included in this review, this was not always the case (e.g. CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014); CLL-IPI V - Bahlo 2016 (SCAN cohort)), and may have led to additional bias to the one assessed by PROBAST.

Applicability of findings to clinical practice and policy

We identified 52 prognostic models for various outcomes. Due to the lack of external validation, we disregarded 40 of these as not (yet) applicable. Nine more prognostic models were validated only one to three times, which we would not consider to be sufficient information to make a judgement of applicability to different settings. The three models included in meta-analysis of the c-statistic offer more information concerning their performance in different populations. The CLL-IPI seems to perform best among the three models in both the development and external validation cohorts. However, due to the relatively small number of external validation studies (N < 10) and serious limitations in reporting, we cannot draw final conclusions regarding the generalisability and usefulness of this model in clinical practice.

The three models that were included in meta-analyses all predict OS. Until 2015, treatment options for individuals with CLL had not changed drastically over time. However, since the introduction of new agents, such as ibrutinib and idelalisib, survival has improved for all risk groups. For individuals with a poor prognosis, there is hope that they will benefit from new and better tailored treatments. Improvements in survival for all risk groups results in systematic underprediction of survival of prognostic models that were developed using data of patients receiving traditional treatment regimens, while the improvement of prognosis especially for highrisk individuals will lead to lowered discriminative ability of a prognostic model. The prognostic models discussed in detail in this review may thus need to be updated, taking this improved baseline risk into account. In contrast, other patient-relevant outcomes, such as time-to-treatment or progression would not be influenced by improved treatment options as long as a watch-and-wait strategy is adopted, and may be an interesting alternative outcome for prediction models. Some of the models were already externally validated for both of these outcomes, and may be included in a future update of this review.

A further limitation of prognostic models in general is that they usually consider only one outcome per model, thereby not considering a balance between different outcomes, such as survival, quality of life, and other outcomes of interest.

CLL has experienced several changes in diagnostic criteria over time (in particular, in 2008 the criterion of having an absolute lymphocyte count (ALC) more than or equal to $5.0\times10^9/L^2$ to a B-cell count more than or equal to $5.0\times10^9/L^2$, changing the categorisation of individuals with clinical monoclonal lymphocytosis (cMBL) or CLL (Cheson 1996; Hallek 2008). We aimed to explore this change in subgroup analysis but did not, due to the low number of included studies with clear division. However, this change is not expected to have an extensive effect on the performance of a model, as prognosis and patient characteristics are comparable in individuals with cMBL and Rai stage I.

Agreements and disagreements with other studies or reviews

To our knowledge, the current review is the first systematic review that aimed to identify all prognostic models developed for untreated individuals with CLL and their corresponding external validation studies.

We are aware of two other systematic reviews that cover one of the prognostic models that we identified, the CLL-IPI (Molica 2018a; Molica 2018b). These systematic reviews aimed at identifying all published studies that have used CLL-IPI to predict the clinical outcome of CLL; one in patients who received chemoimmunotherapy or targeted therapies (Molica 2018a), the other without restriction of therapy (Molica 2018b). There are some differences between these reviews and our systematic review. In Molica 2018a, studies were limited to individuals who received chemoimmunotherapy or targeted therapies and also included studies of relapsed/refractory CLL from conference abstracts. The authors summarised the survival frequencies per CLL-IPI risk group at two years for OS, whereas we pooled OS at five years. Molica 2018b included patients undergoing any treatment, and summarised both OS and TFS at five years. Both these reviews included data from conference proceedings, whereas we did not. Moreover, they did not report measures of discrimination such as the c-statistic (which is pivotal when assessing model performance) and did not assess risk of bias or applicability.

The included studies in Molica 2018b mostly overlapped with our identified studies for OS. The exception was two external validation studies published as abstracts (N = 2, CLL11 study population; Goede 2016 and Ferrer Lores 2016). In contrast, we identified three additional cohorts which had not yet been published at the time of publication of Molica 2018a and Molica 2018b (CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort); CLL-IPI V - Rani 2018 (Indian cohort); CLL-IPI V - Zhu 2018 (Chinese cohort)).

AUTHORS' CONCLUSIONS

Implications for practice

We identified three prognostic models that were validated more than three times for their primary outcome OS: the CLL-IPI, the Barcelona-Brno score, and the MDACC 2007 index score. Of the three models, the CLL-IPI seems to perform best regarding the discrimination between individuals with a good prognosis as compared to individuals with a worse prognosis. However, the number of external validation studies was relatively low to draw definite conclusions. Especially for the very high-risk group, the index underpredicts survival, which may be a result of the selective sample that was used to develop the index.

This review has been developed in a time of rapidly expanding treatment options that may drastically change the prognostic implications for the therapy and longevity of these patients. The external validation cohorts may not be representative anymore of the currently available, improved treatment options. Therefore, the models can be used to provide an approximate classification, but may need testing and updating before being applied to new patients.



Implications for research

This systematic review shows that for CLL, an abundance of studies of prognostic models and scores can be identified. Based on our very inclusive definition of a prognostic score, we found that these studies (including the newer studies) were not performed according to the current standards of prognostic model development. For the future, we recommend authors of model development studies incorporate the most recent checklists and tools, such as CHARMS and TRIPOD, as orientation to avoid common issues. We would also like to emphasise that reporting in more recent papers can benefit from improvements. The minimum reported information should include participant information (recruitment, study design, eligibility criteria, diagnostic criteria used, etc.), predictor assessment, outcome definition, and the relevant performance measures.

Our review also shows that the proportion of external model validation studies as compared to model development studies is very low. Before investing in the development of a new model, we recommend the validation and direct comparison of the existing models in different settings and populations. Most suitable for this aim are well-designed prospective studies that also take into account novel agents and emerging molecular makers. To improve predictive performance, a model can be updated with these markers or tailored to a specific population.

With time, newly identified prognostic factors may be added.

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Skoetz N, Trivella M, Kreuzer KA, Collins G, Köhler N, Wolff M, et al. Prognostic models for chronic lymphocytic leukaemia: an exemplar systematic review and meta-analysis. *Cochrane Database of Systematic Reviews* 2016, Issue 1. Art. No: CD012022. [DOI: 10.1002/14651858.CD012022]

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Baliakas D - Baliakas 2019 (multicentre)

Study characteristics	

General information

Model and type of study

- Baliakas 2019 (development cohort)
- · Development study

Secondary citations

not applicable

Language of publication

• English

Study design

• multicentre, retrospective cohort

Follow-up time

• median 7.1 years (range 0.1 to 33.1 years)

Participants

Number of included persons in the cohort

- 2366 persons*
- *analysis included only 1900 persons with Binet A, divided into M-CLL (n = 1224) and U-CLL (n = 676)

<u>Setting</u>

• 10 European institutions, multicentre study

Recruitment period

· not reported

Age (in years)

• median 64.3 years (range: 22-92 years)

Sex

• 61% male

Stages of disease

• Binet A: 80%; Binet B: 12%; Binet C: 8%

^{*} Indicates the major publication for the study



Baliakas D - Baliakas 2019	(multicentre)	(Continued)
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Treatment

· not reported

Inclusion criteria

• diagnosed CLL, available immunogenetic data

Exclusion criteria

· not reported

Predictors

Number of candidate predictors

- univariable analysis: 15
- multivariable analysis: 15

<u>List of predictors in final model (including cut-points for dichotomised factors)</u>

- M-CLL: TP53 abnormality, trisomy12, subset #2 (IGHV3-21/IGLV3-21 BcR IG)
- · U-CLL: TP53 abnormality, del11q, gender

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• TTFT: evaluated from the diagnostic date until the date of initial treatment; untreated cases were censored at the time of last follow-up.

Additional outcome(s)

NA

Missing data

Participants with any missing data?

- yes
- M-CLL: 306, U-CLL: 288

If yes, how was missing data handled?

• excluded from multivariable analysis if data were missing for one of the factors included in the model

Analysis

Number of participants and number of events (specific time points where reported)

• TTFT: 1900 persons, number of events not reported

Predictor selection method

selection of prognostic factors based on univariable analysis

Statistical method

 decision tree based on binary recursive partitioning and subsequent application of an amalgamation algorithm

Simplification of model?

yes

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)



Baliakas D - Baliakas 2019 (multicentre) (Continued)

Creation of risk groups?

- yes
- M-CLL: 4 risk groups; i) very high risk: Binet C with identical 5- and 10-year TP of 92%; ii) high risk: Binet B, 5y-TP and 10y-TP: 64% and 84%, respectively; iii) intermediate risk: Binet A with one of the following: TP53abn and/or +12 and/or subset #2 membership, 5y-TP and 10y-TP: 40% and 55%, respectively (of note, among 18 non-censored cases with no treatment indication for more than 10 years after diagnosis, 5 (30%) carried TP53abn); and iv) low risk: non-TP53abn/+12/subset #2 Binet A, 5y-TP and 10y-TP: 12% and 25%, respectively
- U-CLL: 5 risk groups; i) very high risk: Binet C with 5- and 10-year TP of 100%; ii) high risk: Binet B, 5y-TP and 10y-TP: 90% and 100%, respectively; iii) intermediate risk: Binet A with one of the following: TP53abn and/or SF3B1mut and/or del(11q), 5y-TP and 10y-TP: 78% and 98%, respectively; iv) low risk: nonTP53abn/SF3B1mut/del(11q) male Binet A, 5y-TP and 10y-TP: 65% and 85%, respectively; and v) very low risk: non-TP53abn/SF3B1mut/del(11q) female Binet A, 5y-TP and 10y-TP: 45% and 65%, respectively

PROBAST: Applicability

Domain 1: Participant selection

high - sample not representative of entire CLL population as only participants with disease stage Binet
 A were analysed

Domain 2: Predictors

low

Domain 3: Outcome

Iow

Notes

Funding and conflict of interest

- "... the Swedish Cancer Society, the Swedish Research Council, the Knut and Alice Wallenberg Foundation, Karolinska Institutet, Stockholm, the Lion's Cancer Research Foundation, Uppsala, the Marcus Borgström Foundation and Selander's Foundation, Uppsala; H2020 "AEGLE, An analytics framework for integrated and personalized healthcare services in Europe" by the EU; H2020 "MEDGENET, Medical Genomics and Epigenomics Network" (No.692298) by the EU; H2020 "CLLassify, Innovative risk assessment for individualizing treatment in chronic lymphocytic leukemia" (No.702714) by the EU; Associazione Italiana per la Ricerca sul Cancro AIRC Investigator grants #20246, and Special Program Molecular Clinical Oncology AIRC 5 per mille #9965; Progetti di Rilevante Interesse Nazionale (PRIN) #2015ZMRFEA, MIUR, Rome,Italy; TRANSCAN-179 NOVEL JTC 2016; project CEITEC 2020 (LQ1601) by MEYS-CZ, project AZV-MH-CZ 15-30015A-4/2015; JCS was funded by Bloodwise (11052, 12036), the Kay Kendall Leukaemia Fund (873), Cancer Research UK (C34999/A18087, ECMC C24563/A15581), Wessex Medical Research and the Bournemouth Leukaemia Fund; Special Program Molecular Clinical Oncology 5 x 1000 No. 10007, Associazione Italiana per la Ricerca sul Cancro Foundation Milan, Italy; Progetto Ricerca Finaliz"
- "KS and PG received research support from Janssen Pharmaceuticals and Gilead Sciences. KS, PG and AH received research support from Novartis SA, Abbvie and Roche Hellas. The other authors declare no conflict of interests in relation to the present study."

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Missing data used as a reason for exclusion: only participants: (quote) "for whom immunogenetic data was available were included in this multicentre retrospective study".
Domain 2: Predictors	Yes	Detailed description of predictor assessment in the appendix
Domain 3: Outcome	Yes	Predefined outcome definition



Baliakas D - Baliakas 2019 (multicentre) (Continued)

Domain 4: Analysis

No

Participants with missing data were excluded from multivariable analyses:

(quote) "we considered only those cases with available data for all the factors included in the model (n = 918 for M-CLL and n = 384 for U-CLL)"; univariable selection of predictors; assumptions of Cox proportional hazards model were

checked.

Overall judgement

No

Baliakas V - Baliakas 2019 (MLL + Scan.)

Study characteristics

General information

Model and type of study

- Baliakas 2019 (MLL + Scan cohort)
- Validation study

Secondary citations

· not applicable

Language of publication

· English

Study design

· retrospective cohort

Follow-up time

not reported

Participants

Number of included persons in the cohort

• 649 persons

Setting

- Munich Leukemia Laboratory (n = 508)
- Scandinavian population-based study (n = 141)

Recruitment period

· not reported

Age (in years)

• median 63.6 years (range: 29-89 years)

Sex

• 62% male

Stages of disease

• Binet A: 100%

Treatment

not reported



Baliakas V - Baliakas 2019	(MLL + Scan.) (Continued) Inclusion criteria
	CLL, disease stage Binet A
	Exclusion criteria
	not reported
Predictors	List of predictors used for validation (and changes between original predictors and predictors in validation study)
	 M-CLL: TP53 abnormality, trisomy12, subset #2 U-CLL: TP53 abnormality and/or SF3B1 mutation, del(11q), gender
	Timing of predictor measurement
	• at diagnosis
Outcome(s)	Primary outcome in study
	 TTFT was evaluated from the diagnostic date until the date of initial treatment, untreated cases wer censored at the time of last follow-up.
	Additional outcome(s)
	• NA
	Outcome in model development
	• TTFT
Missing data	Participants with any missing data?
	• no
	If yes, how was missing data handled?
	not applicable
Analysis	Number of participants and number of events (specific time points where reported)
	time-to-first-treatment: 649 persons, number of events not reported
	Which model was used?
	classification into risk groups
	Was the model updated?
	• no
	Performance measures reported?
	Calibration: not reportedDiscrimination: not reported
	Creation of risk groups?
	• no
PROBAST: Applicability	Domain 1: Participant selection
	unclear - neither eligibility criteria nor recruitment period reported
	Domain 2: Predictors
Prognostic models for newly-	diagnosed chronic lymphocytic leukaemia in adults: a systematic review and meta-analysis (Review)



Baliakas V - Baliakas 2019 (MLL + Scan.) (Continued)

low

Domain 3: Outcome

low

Notes

Funding and conflict of interest

- "... the Swedish Cancer Society, the Swedish Research Council, the Knut and Alice Wallenberg Foundation, Karolinska Institutet, Stockholm, the Lion's Cancer Research Foundation, Uppsala, the Marcus Borgström Foundation and Selander's Foundation, Uppsala; H2020 "AEGLE, An analytics framework for integrated and personalized healthcare services in Europe" by the EU; H2020 "MEDGENET, Medical Genomics and Epigenomics Network" (No.692298) by the EU; H2020 "CLLassify, Innovative risk assessment for individualizing treatment in chronic lymphocytic leukemia" (No.702714) by the EU; Associazione Italiana per la Ricerca sul Cancro AIRC Investigator grants #20246, and Special Program Molecular Clinical Oncology AIRC 5 per mille #9965; Progetti di Rilevante Interesse Nazionale (PRIN) #2015ZMRFEA, MIUR, Rome,Italy; TRANSCAN-179 NOVEL JTC 2016; project CEITEC 2020 (LQ1601) by MEYS-CZ, project AZV-MH-CZ 15-30015A-4/2015; JCS was funded by Bloodwise (11052, 12036), the Kay Kendall Leukaemia Fund (873), Cancer Research UK (C34999/A18087, ECMC C24563/A15581), Wessex Medical Research and the Bournemouth Leukaemia Fund; Special Program Molecular Clinical Oncology 5 x 1000 No. 10007, Associazione Italiana per la Ricerca sul Cancro Foundation Milan, Italy; Progetto Ricerca Finaliz"
- "KS and PG received research support from Janssen Pharmaceuticals and Gilead Sciences. KS, PG and AH received research support from Novartis SA, Abbvie and Roche Hellas. The other authors declare no conflict of interests in relation to the present study."

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Eligibility criteria not reported; not stated whether participants were excluded based on missing values; recruitment period not reported
Domain 2: Predictors	Unclear	Lack of information about predictor assessment, especially since the validation cohort consisted of two separate cohorts which were merged for analysis
Domain 3: Outcome	Unclear	Outcome definition not reported
Domain 4: Analysis	No	Performance measures (calibration or discrimination) not reported
Overall judgement	No	

Barcelona-Brno D - Delgado 2017 (Barcelona cohort)

Study characteristics

General information

Model and type of study

- Delgado 2017 (development cohort (Barcelona))
- · Development study

Secondary citations

not applicable

Language of publication

• English



Barcelona-Brno D - Delgado 2017 (Barcelona cohort) (Continued)

Study design

· retrospective cohort

Follow-up time

• median 99.6 months (range 1-456 months)

Participants

Number of included persons in the cohort

• 524 persons

Setting

· Spain, single-centre study

Recruitment period

· not reported

Age (in years)

· median 62 years (range: 22-93 years)

Sex

• 60% male

Stages of disease

- Rai 0: 62%; Rai I-IV: 38%
- Binet A: 83%; Binet B/C: 17%

Treatment

- 83 received FCR or similar
- 82 received purine analogs without MoAbs
- 86 received alkylating agents
- · 23 received others

Inclusion criteria

• Information at diagnosis included age, clinical stage (Rai and Binet), IgHV mutational status, beta2-microglobulin (B2M), and FISH-detected cytogenetic abnormalities

Exclusion criteria

· not reported

Predictors

Number of candidate predictors

- univariable analysis: 5
- multivariable analysis: 5

List of predictors in final model (including cut-points for dichotomised factors)

• IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberations (del17p, del11q) (binary)

Timing of predictor measurement

• at diagnosis

Outcome(s)

Primary outcome in study



Barcelona-Brno D - Delgado 2017 (Barcelona cohort) (Continued)

• OS: calculated from the date of diagnosis to the date of death or last follow-up

Additional outcome(s)

 TTFT: calculated from the date of diagnosis to the date of first treatment or last follow-up, considering disease-unrelated deaths as competing events

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants and number of events (specific time points where reported)

- overall survival: 524 persons, 212 events
- TTFT: 524 persons, 292 events

Predictor selection method

 aim was the simplification of the CLL-IPI. Therefore, the 5 factors of this model were combined and the most discriminatory combination was chosen. The exact procedure was unclear.

Statistical method

Identification of combination of biomarkers with significant discriminatory value (based on CLL-IPI model)

Simplification of model?

no

Performance measures reported?

- · Calibration: reported upon request
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups (no 11q or 17p and mutated IgHV); (all others); (11q or 17p and unmutated IgHV))
- · at 10 years
- percentage of group without event at time point (low risk: 82%, intermediate risk: 52%, high risk: 27%)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - no clear eligibility criteria and recruitment period reported

Domain 2: Predictors

• low

Domain 3: Outcome

low

Notes

Funding and conflict of interest

- "Red Tematica de Investigacion Cooperativa en Cancer RT, Grant No.: 06/0020/002051 and RD12/0036/0023; Instituto de Salud Carlos III (ISCIII), Grant No.: FISS PI080304; ICGC-CLL Genome Project, Generalitat de Catalunya, Grant No.: 2009SGR1008; "Emili Letang" (T.B.)"
- "The authors declare no conflict of interest."



Barcelona-Brno D - Delgado 2017 (Barcelona cohort) (Continued)

Other comments

- It seemed that the model was built for OS, but also applied to TTFT, no clear description of this process
- The authors have provided additional information (calibration and discrimination, number of events) upon request via email.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Retrospective retrieval of individuals with available information for their model - individuals without the necessary data may have been left out. Unclear recruitment period
Domain 2: Predictors	Yes	No explicit statement on predictor measurement, however, there are indications that in this single-centre study, lab procedures remained similar. This was a retrospective cohort, possibility to look forward
Domain 3: Outcome	Yes	No clear description of outcome assessment. We assumed that assessment of the objective outcome OS was similar within this single-centre study and therefore did not rate as high risk (no information on e.g. registry, frequency of follow-up)
Domain 4: Analysis	No	Dichotomisation of predictors. Although the choice of predictors was based on a previous model (CLL-IPI), no formal factor selection procedure or comparison of tested factor combinations were reported. One point assigned per factor, no formally established factor weights
Overall judgement	No	One point assigned per factor, no formally established factor weights

arcelona-Brno V - Delgado 2017 (Brno cohort) Study characteristics		
General information	Model and type of study	
	Delgado 2017 (Brno cohort, Czech Republic)Validation study	
	<u>Secondary citations</u>	
	not applicable	
	Language of publication	
	• English	
	Study design	
	not reported	
	Follow-up time	
	median 59 months (range 3-330 months)	
Participants	Number of included persons in the cohort	



Barcelona-Brno V - Delgado 2017 (Brno cohort) (Continued)

· 417 persons

Setting

• Czech Republic (Brno), single-centre study

Recruitment period

· not reported

Age (in years)

• median 62 years (range: 32-85 years)

Sex

• 66% male

Stages of disease

- Rai 0: 41%; Rai I-II: 43%; Rai III-IV: 16%
- Binet A: 72%; Binet B: 12%; Binet C: 16%

Treatment

- 125 received FCR or similar
- 41 received purine analogs without MoAbs
- 53 received alkylating agents
- · 57 received others

Inclusion criteria

not reported

Exclusion criteria

not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberations (del17p, del11q) (binary)
- no differences between development and validation predictors

Timing of predictor measurement

• at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: OS was calculated from the date of diagnosis to the date of death or last follow-up.

Additional outcome(s)

• TTFT: calculated from the date of diagnosis to the date of first treatment or last follow-up, considering disease-unrelated deaths as competing events

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no



Barcelona-Brno V - Delgado 2017 (Brno cohort) (Continued)

If yes, how was missing data handled?

not applicable

Analysis

Number of participants and number of events (specific time points where reported)

- OS: 417 persons, 158 events
- TTFT: 417 persons, 276 events

Which model was used?

· original model

Was the model updated?

no

Performance measures reported?

- Calibration: provided upon request (calibration plot)
- · Discrimination: provided upon request (c-statistic)

Creation of risk groups?

- yes, 3 risk groups (no 11q or 17p and mutated IgHV); (all others); (11q or 17p and unmutated IgHV)
- at 5 years
- percentage of group without event at time point (low risk: 88.4% (83.0 to 94.1%), intermediate risk: 72.5% (65.4 to 80.4%), high risk: 53.9% (44.2 to 65.8%))

PROBAST: Applicability

Domain 1: Participant selection

· unclear - recruitment period and eligibility criteria unclear

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding and conflict of interest

- "Ministry of Health of the Czech Republic, Grant No.: AZV 15-31834A/ 2015 and AZV 15-30015A/2015; Ministry of Education, Youth and Sports of the Czech Republic project NPUII - CEITEC 2020, Grant No.: LQ1601."
- "The authors declare no conflict of interest."

Other comments

• The authors have provided additional information (calibration and discrimination, number of events) upon request via email.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se-	Unclear	No recruitment period and study design reported
lection		No clear inclusion and exclusion criteria (we do not know if missing values have been part of the exclusion criteria)



Barcelona-Brno V - Delgado 2017 (Brno cohort) (Continued)

Domain 2: Predictors	Yes	Not explicitly stated, but predictors probably assessed in a similar way
Domain 3: Outcome	Yes	No clear description of outcome assessment. We assumed that assessment of the objective outcome OS was similar within this single-centre study and therefore did not rate as high risk (no information on e.g. registry, frequency of follow-up)
Domain 4: Analysis	Unclear	No information on missing values; information on calibration and discrimination was provided upon request

Barcelona-Brno V - Gentile 2017 (Italian & Mayo)

Study	charac	teristics

General information

Model and type of study

- Gentile 2017 (Italian multicentre and Mayo clinic cohort)
- Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

• median 82 months (range 3-330 months)

Participants

Number of included persons in the cohort

• 1299 persons

Setting

• Italy, USA, multicentre study

Recruitment period

• 1985 - 2015

Age (in years)

• median 63 years (range: 27-92 years)

Sex

• 61.3% male

Stages of disease

- Rai 0: 57.9%; Rai I: 28.6%; Rai II: 8.6%; Rai III: 1.5%; Rai IV: 3.5%
- Binet A/B/C: not reported

Treatment



Barcelona-Brno V - Gen	tile 2017 (Italian & Mayo) (Continued) • 194 received chemotherapy			
	316 received chemoimmunotherapy			
	Inclusion criteria			
	newly-diagnosed CLL persons			
	Exclusion criteria			
	not reported			
Predictors	<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>			
	 IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberations (del17p, del11q) (binary) no differences between development and validation predictors 			
	Timing of predictor measurement			
	at diagnosis			
Outcome(s)	Primary outcome in study			
	• overall survival: for OS, the time interval was measured from the day of CLL diagnosis until death from all causes or last follow-up.			
	Additional outcome(s)			
	TTFT: measured as the day of CLL diagnosis until the start of therapy or last follow-up			
	Outcome in model development			
	overall survival			
Missing data	Participants with any missing data?			
	• no			
	If yes, how was missing data handled?			
	not applicable			
Analysis	Number of participants and number of events (specific time points where reported)			
	OS: 1299 persons, 283 eventsTTFT: 1299 persons, 510 events			
	Which model was used?			
	original model			
	Was the model updated?			
	• no			
	Performance measures reported?			
	Calibration: reported (explained variation)Discrimination: reported (c-statistic)			
	Creation of risk groups?			

• yes, 3 risk groups (no 11q or 17p and mutated IgHV); (all others); (11q or 17p and unmutated IgHV)



Barcelona-Brno V - Gentile 2017 (Italian & Mayo) (Continued)

- · at 5 years
- percentage of group without event at time point (low risk: 92.2%, intermediate risk: 83.6%, high risk: 68.2%)

PROBAST: Applicability

Domain 1: Participant selection

 unclear - recruitment period spanned a broad time period, wherein CLL definitions and treatment options have changed

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding and conflict of interest

- · Funding not reported
- "The authors have no conflict of interest to disclose."

Other comments

- This publication referred to the same cohort as in Gentile 2016
- The authors provided no additional information on calibration upon request.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used. Many missing values due to retrospective design, rated in domain 4
Domain 2: Predictors	Yes	Quote: "IgHV mutation analysis and FISH were performed at the reference laboratory of each participating center. The IgHV mutation status was tested on tumour DNA collected at diagnosis, and was assessed according to the ERIC guidelines."
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment. We assumed that assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	> 70% of individuals were excluded based on missing values (quote: "because of the absence of the required laboratory data in the cohort initially considered. However, the individuals included were representative of the whole cohort and were similar for age, sex, and Rai stage distribution").
Overall judgement	No	

Barcelona-Brno V - Molica 2017 (O-CLL1-GISL)

Study characteristics

General information

Model and type of study

- Molica 2017 (O-CLL1-GISL)
- Validation study



Barcelona-Brno V - Molica 2017 (O-CLL1-GISL) (Continued)

Secondary citations

· Delgado 2017

Language of publication

• English

Study design

· prospective cohort

Follow-up time

• median 42 months (range 1 - 82 months)

Participants

Number of included persons in the cohort

· 337 persons

Setting

· Italy, multicentre study

Recruitment period

· 2007 - not reported

Age (in years)

• median 61 years (range: 33 - 70 years)

Sex

• 57.2% male

Stages of disease

- Rai 0: 77.8%; Rai I-II: 22.2%
- Binet A: 100%; Binet B/C: not applicable

Treatment

· not reported

Inclusion criteria

Newly-diagnosed individuals with CLL from several Italian Institutions who were seen within 12 months of diagnosis (confirmed by the biological review committee according to flow cytometry analysis (positive clusters of differentiation antigen 5 (CD5), 19 (CD19), 23 (CD23)); established diagnosis of B-CLL by NCI criteria, performed by local haematologist; age above 18 years and below 70 years; Binet stage A; NCI watch-and-wait policy

Exclusion criteria

 diagnosis more than 12 months ago, aged above 70 years, patients with leukaemic phase of lymphoproliferative disorders of B origin CD5- and/or CD23- according to flow cytometry analysis, Binet B or C

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberrations (del17p, del11q) (binary)
- TP53 and del17p were used as composite predictor in place of del17p only.

Timing of predictor measurement



Barcelona-Brno V - Molica 2017 (O-CLL1-GISL) (Continued)

• at study entry, within 12 months of diagnosis

Outcome(s)

Primary outcome in study

 time-to-first-treatment: the primary endpoint, TTFT, was defined as the interval between the date of registration and the date of initiation of first CLL treatment. (Molica 2016)

Additional outcome(s)

· not applicable

Outcome in model development

overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 337 persons, 91 events

Which model was used?

· original model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- · Discrimination: reported (AUC)

Creation of risk groups?

no

PROBAST: Applicability

Not applicable; outcome did not match primary outcome of the model

Notes

Funding & conflict of interest

- "This work was supported by funding from Associazione Italiana per la Ricerca sul Cancro (AIRC 5xmille grant 9980, IG10492 to MF and FM and IG10136 to AN). We thank AIL Cosenza-Fondazione Amelia Scorza' onlus, Cosenza, Italy, and Brigida Gulino for precious secretarial assistance." (Gentile 2016)
- "The authors have no conflict of interest to disclose."

Other comments

- This publication validated the Barcelona-Brno score in the same population as the validation cohort in the development study.
- The outcome did not correspond to the primary outcome of the model, and was therefore not included for analysis and PROBAST rating.



Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.)

Study characteristics

General information

Model and type of study

- Munoz-Novas 2018 (Spanish cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

• median 68 months (range 3 - 277 months)

Participants

Number of included persons in the cohort

· 696 persons

Setting

· Spain, multicentre study

Recruitment period

• 1989 - 2013

Age (in years)

• median 65.7 years (IQR: 55.2 - 73.5)

Sex

• 62.8% male

Stages of disease

- Rai 0: 58.9%; Rai I-IV: 41.1%
- Binet A/B/C: not reported

Treatment

- 137 received treatment
- unclear which treatment

Inclusion criteria

• all persons diagnosed with CLL

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.) (Continued)

- IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberations (del17p, del11q) (binary)
- no differences between development and validation predictors

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

 overall survival: Overall survival (OS) was calculated from the time of diagnosis to death or last follow-up.

Additional outcome(s)

• time-to-first-treatment: Time-to-first therapy (TTFT) was defined from the date of diagnosis to first treatment or last follow-up.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

• yes (438 persons)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 258 persons, 47 events
- time-to-first-treatment: 258 persons, 113 events

Which model was used?

· original model

Was the model updated?

no

Performance measures reported?

- · Calibration: not reported
- · Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups (no 11q or 17p and mutated IgHV); (all others); (11q or 17p and unmutated IgHV)
- at 5 years
- percentage of group without event at time point (low risk: 90.7%, intermediate risk: 81.4%, high risk: 66.2%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome



Barcelona-Brno V - Muñoz-Novas 2018 (Spanish coh.) (Continued)

low

Notes

Funding & conflict of interest

- · Funding not reported
- "The authors declare that there are no conflicts of interest regarding the publication of this article."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Although individuals with incomplete data were excluded, there was a comparison and the sample seemed representative of the cohort.
Domain 2: Predictors	Yes	The type of biomarkers used seem to be standard, and although there was no explicit information we think that the methods were consistent across studies and models.
Domain 3: Outcome	Yes	Objective standard outcome - no clear description of outcome assessment - we assumed that assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	Consecutive sampling of individuals in hospital routine, therefore around 2/3 of individuals with missing data for this model validation were excluded; no calibration reported.
Overall judgement	No	

Barcelona-Brno V - Rani 2018 (Indian cohort)

Study characteristics

General information

Model and type of study

- Rani 2018 (Indian cohort)
- Validation study

Secondary citations

• not applicable

Language of publication

• English

Study design

unclear

Follow-up time

• median 40.5 months (range 1 - 215 months)

Participants

Number of included persons in the cohort



Barcelona-Brno V - Rani 2018 (Indian cohort) (Continued)

198 persons

Setting

• India, unclear if single or multicentre study

Recruitment period

· not reported

Age (in years)

• median 60 years (range: not reported)

Sex

• 77% male

Stages of disease

- Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%
- Binet A/B/C: not reported

Treatment

- 62 received chlorambucil-based therapy
- · 56 received rituximab-based therapy
- 20 received other therapies

Inclusion criteria

· treatment-naive CLL persons according to Hallek 2008

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberations (del17p, del11q) (binary)
- no differences between development and validation predictors

Timing of predictor measurement

not reported

Outcome(s)

Primary outcome in study

• overall survival: OS was defined as the time from the date of diagnosis to date of death or date of last follow-up.

Additional outcome(s)

• time-to-first-treatment: TTFT was defined as the time from the date of diagnosis to date of commencement of first therapy.

Outcome in model development

overall survival

Missing data

Participants with any missing data?

no



Barcelona-Brno V - Rani 2018 (Indian cohort) (Continued)

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 198 persons, 86 events
- time-to-first-treatment: 140 persons, 89 events

Which model was used?

· original model

Was the model updated?

no

Performance measures reported?

- · Calibration: not reported
- · Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups (no 11q or 17p and mutated IgHV); (all others); (11q or 17p and unmutated IgHV)
- median survival (low risk: 138 years, intermediate risk: 63 years, high risk: 56 years)

PROBAST: Applicability

Domain 1: Participant selection

· unclear - recruitment period and eligibility criteria unclear

Domain 2: Predictors

• unclear - no information on timing of predictor because the study design is unclear

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- "... Department of Biotechnology (BT/PR11106/GBD/27/145/2008, BT/PR15438/MED/30/606/2011 and T/PR8680/AGR/36/754/2013), Ministry of Science and Technology, GOI, and All India Institute of Medical Sciences, New Delhi (8-60/A060/ 2011/RS) to RG to carry out this work."
- "The authors declare that they have no conflict of interest."

Other comments

none

Authors' judgement	Support for judgement
Unclear	Inclusion and exclusion criteria unclear; it seemed to us that only individuals with complete information on all predictors were included.
Yes	Not explicitly stated, but predictors probably assessed in a similar way (single-centre)
	Unclear



Barcelona-Brno V - Rani 2018 (Indian cohort) (Continued)

Domain 3: Outcome	Unclear	Observation time to observe survival was short.
Domain 4: Analysis	No	Low number of events, no information on handling of missing values; calibration was not reported.
Overall judgement	No	

Barcelona-Brno V - Reda 2017 (Milan cohort)

Study cho	aracte	ristics
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General information

Model and type of study

- Reda 2017 (Milan cohort)
- Validation study

Secondary citations

• not applicable

Language of publication

English

Study design

· retrospective cohort

Follow-up time

• median 144 months (range 0 - 360 months)

Participants

Number of included persons in the cohort

• 698 persons

Setting

• Italy, single-centre study

Recruitment period

• 1983 - 2016

Age (in years)

• median 65 years (range: 32 - 91 years)

<u>Sex</u>

• 57% male

Stages of disease

- Rai 0-I: 74%; Rai II-IV: not reported
- Binet A/B/C: not reported

Treatment

- 190 received Chlorambucil (CHL)
- 108 received fludarabinecyclophosphamide-rituximab (FCR)



Barcelona-Brno V - Reda 2017 (Milan cohort) (Continued)

- 12 received of atumum abbendamustine (O-Benda)
- 44 received bendamustine-rituximab (BR)
- 65 received alemtuzumab
- 29 received Ibrutinib or Idelalisib

Inclusion criteria

· all patients diagnosed with CLL, reclassified by 2008 criteria

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- IgHV mutational status (continuous: categorical/binary transformation; binary, cut-off not given but usually at 98%); FISH-detected genetic abberations (del17p, del11q) (binary)
- no differences between development and validation predictors

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: overall survival (OS) was calculated from CLL diagnosis to death.

Additional outcome(s)

 time-to-first-treatment: Time-To-First-Treatment (TTFT) was evaluated from the time of diagnosis to firstline therapy start.

Outcome in model development

overall survival

Missing data

Participants with any missing data?

• yes (369 participants)

If yes, how was missing data handled?

unclear

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 329 persons, number of events not reported
- time-to-first-treatment: 329 persons, number of events not reported

Which model was used?

· original model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: not reported

Creation of risk groups?



Barcelona-Brno V - Reda 2017 (Milan cohort) (Continued)

- yes, 3 risk groups (no 11q or 17p and mutated IgHV); (all others); (11q or 17p and unmutated IgHV)
- at 15 years
- percentage of group without event at time point (low risk: 80%, intermediate risk: 50%, high risk: 30%)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - no clear information on eligibility criteria and study design

Domain 2: Predictors

• unclear - no information on the timing of the predictor

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- · Funding not reported
- "Authors have no affiliations that they consider to be relevant and important with any organization
 that to any author's knowledge has a direct interest, particularly a financial interest, in the subject
 matter discussed."

Other comments

• The authors did not reply to our request via email for additional information.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	No eligibility criteria, unclear study design
Domain 2: Predictors	Yes	Not explicitly stated, but predictors probably assessed in a similar way
Domain 3: Outcome	Yes	No clear description of outcome assessment - we assumed that assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	No information on number of events; no performance measures reported; patients with missing data were left out from analysis.
Overall judgement	No	

CLL-IPI D - Bahlo 2016 (development cohort)

Study characteristics

General information

Model and type of study

- Bahlo 2016 (development cohort)
- · Development study

Secondary citations

· not applicable

Language of publication



English

Study design

• combined data (e.g. IPD or combination of cohorts); data from 8 phase-three RCTs

Follow-up time

• median 79.9 months, IQR: 79.9 - 101.4

Participants

Number of included persons in the cohort

• 3472 persons

Setting

- · multicentre study
- France, Germany, Poland, UK, USA, and possibly more

Recruitment period

1997 - 2009

Age (in years)

• median 61 years (range: 27 - 86 years)

Sex

• 70% male

Stages of disease

- Rai 0: 14%; Rai I/II: 50%; Rai III/IV: 36%
- Binet A: 32%; Binet B: 41%; Binet C: 27%

Treatment

- 690 received fluradabine (F)
- 1017 received F + cyclophosphamide (C)
- 113 received C + cladribine
- 475 received FC + rituximab
- 81 received FC + alemtuzumab
- 479 received chlorambucil

Inclusion criteria

untreated CLL - possibly more exclusion criteria in the individual RCTs; FR: between the ages of 18 and 65 years; absence of del17p, life expectancy longer than 6 months; DE - CLL4: 18 and 65 years; life expectancy more than 6 months; Binet stage C and stage B if they had rapid disease progression or symptoms as evidenced by enlarged lymph nodes and organs or if they had severe B symptoms, stage A with severe B-symptoms; DE - CLL5: 65 and 80 years; life expectancy more than 6 months; Binet stage C and in stage B or A if they had rapid disease progression (lymphocyte doubling time 3 months) or symptoms from enlarged lymph nodes and organs, or if they had severe B-symptoms; PL: more than 18 years, progressive disease; UK: stages B, C, and A-progressive who needed treatment; USA: 18 years or older with a diagnosis of progressive CLL

Exclusion criteria

• FR: comorbidities (CIRS more than 5, ECOG more than 2; autoimmune cytopenia, HIV, HBV, HCV, active second malignancy, transformation to aggressive B-cell malignancy, abnormal renal function, bilirubin, transaminase more than 2 times the upper limit; DE - CLL 4: ECOG more than 3; organ dysfunction, concomitant or previous neoplasms, autoimmune haemolytic anaemia or thrombocytopenia; DE - CLL5: ECOG more than 3; organ dysfunction, concomitant or previous neoplasias, autoimmune



haemolytic anaemia, thrombocytopenia; DE - CLL8: ECOG more than 2; CIRS more than 5; autoimmune cytopenia, active second disease; PL: poor performance status (WHO grade 4), autoimmune haemolytic anaemia (AIHA), autoimmune thrombocytopenia, active infections, abnormal liver or renal function, Richter's syndrome, or concomitant neoplasm; USA: second malignancy, serum creatinine more than 2 mg/dL; performance status more than 2; bilirubin more than 1 mg/dL; autoimmune haemolytic anaemia, thrombocytopenia; history of steroid treatment; pregnancy and lactating women; active infection

Predictors

Number of candidate predictors

- univariable analysis: 27
- multivariable analysis: 25

List of predictors in final model (including cut-points for dichotomised factors)

TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L based on literature); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off: 65 years)

Timing of predictor measurement

· before the start of first-line treatment

Outcome(s)

Primary outcome in study

• overall survival: overall survival was calculated from study entry to date of death.

Additional outcome(s)

· not applicable

Missing data

Participants with any missing data?

· yes (1673 participants)

If yes, how was missing data handled?

· complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

 overall survival: 1214 persons, 462 events for training set; 585 persons, 243 events for internal validation set

Predictor selection method

univariable analysis, during multivariable modelling: a stepwise modelling based on categories of missing values: step 1: N = 1908, age, gender, Binet, LDH, haemoglobin [ex: WBC, HGB, LYM]; step 2: N = 1563 (585 events), age, gender, clinical stage (Rai OR Binet), LDH, haemoglobin, ECOG, hierarchical type (del17p, del11q); step 3: N = 1192 (452 events); age, clinical stage (Rai or Binet), IgHV, B2M, hierarchical type (del17p); step 4: more MV analyses (table S11 P value or CI)

Statistical method

Cox proportional hazard model

Simplification of model?

yes

Performance measures reported?

· Calibration: not reported



· Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- at 5 years
- percentage of group without event at time point (low risk: 93.2%, intermediate risk: 79.3%, high risk: 63.3%, very high risk: 23.3%)

PROBAST: Applicability

Domain 1: Participant selection

 high - most individuals had a previous treatment indication, in total only 17.8% of individuals were on watch-and-wait strategy at enrolment, therefore the population was rather selective.

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "José Carreras Leukaemia Foundation"
- KB: BMBF grant. BFE: honorarium for advisory boards, honorarium and/or scientific grants from Janssen, Gilead, Mundipharma, GlaxoSmithKline (GSK) and Roche. CG: personal fees from Roche, Janssen, Gilead, Celgene, Novartis, and AbbVie, outside the submitted work. MG: National Cancer Institute during the conduct of the study; grants, personal fees, and non-financial support from Pharmacyclics and Acerta, outside the submitted work. TDS: grants from Genentech, Pharmacyclics Janssen, GSK, Celgene, Cephalon, Hospira, and Polyphenon E International, outside the submitted work. SS: grants, personal fees, and other from AbbVie, Amgen, Boehringer Ingelheim, Celgene, Genentech, Genzyme, Gilead, GSK, Janssen, Mundipharma, Novartis, Pharmacyclics, Hoffmann La-Roche, and Sanofi, during the conduct of the study. "JBa, NK, MAB, JBy, KGC, HD, ME, SL, DN, DO, TR, RR, and MH declare no competing interests." "The funder of the study had no role in the study design, data collection, data analysis, data interpretation, or writing of the report."

Other comments

The authors provided additional information upon request (cohort information, measures of calibration).

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used, missing values were excluded from analysis. This was rated in domain 4.
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way, according to RCT protocols.
Domain 3: Outcome	Yes	Objective standard outcome, follow-up in the context of each individual RCT
		Median observation time seemed on the lower limit (around 5 years), but reasonable for patients with treatment indication.
Domain 4: Analysis	No	It was unclear for which model the performance was assessed (score or formula). Predictors were categorised and dichotomised. Patients with missing values were dropped from the analysis - exclusion of approximately half of all patients. Univariable analysis was used for predictor selection.



Upon request, the authors provided information on the calibration of their model.

Overall judgement

No

CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014)

Study characteristics

General information

Model and type of study

- Bahlo 2016 (Mayo clinic 2001 2014)
- · Validation study

Secondary citations

• Gentile 2016; Molica 2016

Language of publication

· English

Study design

· prospective cohort

Follow-up time

• median 63.2 months, IQR 30.2 to 91.8

Participants

Number of included persons in the cohort

· 838 persons

Setting

· USA, single-centre study

Recruitment period

• 2001 - 2014

Age (in years)

- ≤ 65 years: 533 persons
- > 65 years: 305 persons

Sex

• 67.8% male

Stages of disease

- Rai 0: 57.4%; Rai I-II: 38.5%; Rai III-IV: 4.1%
- Binet A/B/C: not reported

<u>Treatment</u>

- 163 received purine analogue chemotherapy
- 5 received purine analogue monotherapy or combination without monoclonal antibody
- · 37 received alkylator-based chemoimmunotherapy



CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014) (Continued)

- 11 received alkylator monotherapy or combination without monoclonal antibody
- 97 received antibody only
- · 7 received RTK inhibitors
- 6 received other treatments

Inclusion criteria

complete baseline data

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); Age (continuous: categorical/binary transformation; cut-off 65 years)
- missing data for TP53, therefore replaced by del(17p)

Timing of predictor measurement

· not reported, presumably at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: OS calculated from diagnosis to death.

Additional outcome(s)

· not applicable

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• overall survival: 838 persons, 144 events

Which model was used?

· simplified model

Was the model updated?

nc

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?



CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014) (Continued)

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- at 5 years
- percentage of group without event at time point (low risk: 96.6%, intermediate risk: 92%, high risk: 68.5%, very high risk: 21.2%)

PROBAST: Applicability

Domain 1: Participant selection

 unclear - insufficient information on CLL diagnostic criteria and inclusion criteria (i.e. availability of baseline data).

Domain 2: Predictors

• unclear - predictor TP53 not available and replaced by a proxy, del(17p)

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "José Carreras Leukaemia Foundation"
- KB: BMBF grant. BFE: honorarium for advisory boards, honorarium and/or scientific grants from Janssen, Gilead, Mundipharma, GlaxoSmithKline (GSK) and Roche. CG: personal fees from Roche, Janssen, Gilead, Celgene, Novartis, and AbbVie, outside the submitted work. MG: National Cancer Institute during the conduct of the study; grants, personal fees, and non-financial support from Pharmacyclics and Acerta, outside the submitted work. TDS: grants from Genentech, Pharmacyclics Janssen, GSK, Celgene, Cephalon, Hospira, and Polyphenon E International, outside the submitted work. SS: grants, personal fees, and other from AbbVie, Amgen, Boehringer Ingelheim, Celgene, Genentech, Genzyme, Gilead, GSK, Janssen, Mundipharma, Novartis, Pharmacyclics, Hoffmann La-Roche, and Sanofi, during the conduct of the study. "JBa, NK, MAB, JBy, KGC, HD, ME, SL, DN, DO, TR, RR, and MH declare no competing interests." "The funder of the study had no role in the study design, data collection, data analysis, data interpretation, or writing of the report."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	No information on inclusion criteria, definition of CLL and if/how many patients were excluded due to missing baseline data
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way (single-centre)
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment - we assumed that assessment of the objective outcome OS was similar and therefore did not rate as high risk (no info on e.g. registry, frequency of follow-up etc.)
Domain 4: Analysis	Yes	Number of events sufficient
		Patients with missing values possibly dropped at enrolment, not clearly stated: 'There were no missing values for the MAYO cohort.'. This was rated in domain 1.
		Upon request, the authors provided information on the calibration of their model.



CLL-IPI V - Bahlo 2016 (Mayo clinic 2001-2014) (Continued)

Overall judgement

No

CLL-IPI V - Bahlo 2016 (SCAN cohort)

Study characteristics

General information

Model and type of study

- Bahlo 2016 (SCAN cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

 other: cohort of CLL patients within a so-called case-control study, which was actually a register; uncertain whether prospective or retrospective study design

Follow-up time

• median 151 months, IQR 124.8 to 163.0

Participants

Number of included persons in the cohort

· 416 persons

Setting

• Denmark and Sweden, multicentre study

Recruitment period

• 1999 - 2002

Age (in years)

- ≤ 65 years: 242 persons
- > 65 years: 174 persons

Sex

• 62.7% male

Stages of disease

- Rai: not reported
- Binet A: 78.1%; Binet B: 16.6%; Binet C: 5.3%

<u>Treatment</u>

· not reported

Inclusion criteria

• complete baseline data



CLL-IPIV - Bahlo 2016 (SCAN cohort) (Continued)

Exclusion criteria

The source population for the SCALE study was restricted to subjects with sufficient knowledge of the
Danish or Swedish language to answer questions in a telephone interview and without a history of
organ transplantation, human immunodeficiency virus infection, or other hematopoietic malignancy
(Smedby 2005).

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- 4% of ß2-microglobulin imputed (multiple imputation)

Timing of predictor measurement

· not reported, presumably at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: OS was calculated from diagnosis to death.

Additional outcome(s)

· not applicable

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

yes (17 participants)

If yes, how was missing data handled?

· multiple imputation

Analysis

Number of participants & number of events (specific time points where reported)

• overall survival: 416 persons, 215 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- at 5 years
- percentage of group without event at time point (low risk: 92.1%, intermediate risk: 75%, high risk: 64.3%, very high risk: 28.6%)



CLL-IPIV - Bahlo 2016 (SCAN cohort) (Continued)

PROBAST: Applicability

Domain 1: Participant selection

unclear - the SCALE study database encompassed the entire population in Denmark and Sweden in
a specified time frame. For this model validation, only the CLL cohort was relevant. We did not know
exactly how many participants were excluded due to missing information. No eligibility criteria were
defined. We were unsure if the population can reflect the general CLL population.

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "José Carreras Leukaemia Foundation"
- KB: BMBF grant. BFE: honorarium for advisory boards, honorarium and/or scientific grants from Janssen, Gilead, Mundipharma, GlaxoSmithKline (GSK) and Roche. CG: personal fees from Roche, Janssen, Gilead, Celgene, Novartis, and AbbVie, outside the submitted work. MG: National Cancer Institute during the conduct of the study; grants, personal fees, and non-financial support from Pharmacyclics and Acerta, outside the submitted work. TDS: grants from Genentech, Pharmacyclics Janssen, GSK, Celgene, Cephalon, Hospira, and Polyphenon E International, outside the submitted work. SS: grants, personal fees, and other from AbbVie, Amgen, Boehringer Ingelheim, Celgene, Genentech, Genzyme, Gilead, GSK, Janssen, Mundipharma, Novartis, Pharmacyclics, Hoffmann La-Roche, and Sanofi, during the conduct of the study. "JBa, NK, MAB, JBy, KGC, HD, ME, SL, DN, DO, TR, RR, and MH declare no competing interests." "The funder of the study had no role in the study design, data collection, data analysis, data interpretation, or writing of the report."

Other comments

• Upon request, the authors provided information on the calibration of their model.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	The SCALE study database encompassed the entire population in Denmark and Sweden in a specified time frame. For this model validation, only the CLL cohort was relevant. We did not know exactly how many participants were excluded. No eligibility criteria were defined.
Domain 2: Predictors	Yes	Patient data collection between 1999 and 2002; we assumed that within this time frame, predictor assessment remained relatively homogenous.
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment in this publication. However, this was a national cohort study, therefore we assumed standardised outcome assessment.
Domain 4: Analysis	Yes	Sufficient number of events
		Missing data was handled appropriately: 'we analysed missing values of the SCAN cohort using Little's MCAR test and imputed these using linear regression.'
		Upon request, the authors provided information on the calibration of their model.



CLL-IPIV - Da Cunha-Bang 2016 (Danish cohort)

Study characteristics

General information

Model and type of study

- Da Cunha-Bang 2016 (Danish National CLL Registry)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

· prospective cohort

Follow-up time

• median 38.4 months, range not reported

Participants

Number of included persons in the cohort

3023 persons

Setting

• Denmark, multicentre study

Recruitment period

• 2008 - 2015

Age (in years)

• median 69 years (range: not reported)

Sex

• 60% male

Stages of disease

- Rai: not reported
- Binet A: 80%; Binet B/C: 20%

Treatment

- 295 received treatment
- · unclear which treatment

Inclusion criteria

• all patients from national registry

Exclusion criteria

· missing data

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



CLL-IPI V - Da Cunha-Bang 2016 (Danish cohort) (Continued)

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- no differences between development and validation predictors reported

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• progression-free survival: TTE (time-to-event), event defined as treatment or death.

Additional outcome(s)

· overall survival: OS defined as time to death

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- progression-free survival: 1514 persons, 544 events
- overall survival: 1514 persons, 249 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- · Discrimination: not reported

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- no time point and numerical results reported, see figure 1 instead

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

• unclear - predictors not described, therefore it was unclear if TP53 or a proxy was used

Domain 3: Outcome

low



CLL-IPI V - Da Cunha-Bang 2016 (Danish cohort) (Continued)

Notes

Funding & conflict of interest

- · Funding not reported
- "During the study, C.U.N. received grants from the Danish Cancer Society, consultancy fees (from Janssen, Roche, Abbvie, and Gilead), and grants (from Novartis and Roche) outside the submitted work and is the principal investigator for clinical trials sponsored by Roche. The remaining authors declare no competing financial interests."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Individuals with missing data previously excluded; comparison of included and excluded individuals showed similar baseline characteristics.
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way. Relatively recent and prospectively followed cohort
Domain 3: Outcome	Unclear	For mortality, the observation time was too short (our clinician recommended observation time to exceed 5 years, median observation time here: 3.2 years)
Domain 4: Analysis	Unclear	Patients with missing data previously excluded, although authors stated that they were comparable. Rated in domain 1.
		No information on model performance measures, however, a validation in our sense was not planned.

CLL-IPI V - Delgado 2017 (Barcelona cohort)

Study	charac	teristics

General information

Model and type of study

- Delgado 2017 (Barcelona hospital cohort)
- Validation study

Secondary citations

not applicable

Language of publication

· English

Study design

· retrospective cohort

Follow-up time

• median 99.6 months, range not reported

Participants

Number of included persons in the cohort



CLL-IPI V - Delgado 2017 (Barcelona cohort) (Continued)

524 persons

Setting

• Spain (Barcelona), single-centre study

Recruitment period

· not reported

Age (in years)

• median 62 years (range: 22 - 93 years)

Sex

• 60% male

Stages of disease

- Rai 0: 62%; Rai I-IV: 38%
- Binet A: 83%; Binet or C: 17%

Treatment

- 83 received FCR or similar
- 82 received purine analogs w/o MoAbs
- · 86 received alkylating agents
- · 23 received others

Inclusion criteria

• Availability of information on age, clinical stage, IgHV status, ß2m, FISH

Exclusion criteria

not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- no changes

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: OS was calculated from the date of diagnosis to the date of death or last follow-up.

Additional outcome(s)

not applicable

Outcome in model development

overall survival

Missing data

Participants with any missing data?



CLL-IPI V - Delgado 2017 (Barcelona cohort) (Continued)

n

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

· overall survival: 524 persons, number of events not reported

Which model was used?

simplified model

Was the model updated?

no

Performance measures reported?

- · Calibration: not reported
- · Discrimination: not reported

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10), using Rai or Binet)
- at 5 years
- percentage of group without event at time point (low risk: 95.7%, intermediate risk: 87.1%, high risk: 63.5%, very high risk: 51.9%)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - no clear eligibility criteria and recruitment period reported

Domain 2: Predictors

• low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "Red Tematica de Investigacion Cooperativa en Cancer RT, Grant No.: 06/0020/002051 and RD12/0036/0023; Instituto de Salud Carlos III (ISCIII), Grant No.: FISS PI080304; ICGC-CLL Genome Project, Generalitat de Catalunya, Grant No.: 2009SGR1008; "Emili Letang" (T.B.)."
- "The authors declare no conflict of interest."

Other comments

 Upon request, the authors provided additional information on the discrimination and calibration of the CLL-IPI model in their cohort.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Retrospective retrieval of persons with available information for their model - individuals without the necessary data may have been left out. Unclear recruitment period



CLL-IPI V - Delgado 2017 (Barcelona cohort) ((Continued)
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way (single-centre study)
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment in this publication. We assumed standardised outcome assessment (single-centre study).
Domain 4: Analysis	Yes	Sufficient number of events.
		No missing values, however, possibly an exclusion reason. This was rated in domain 1.
		Upon request, the authors provided information on the calibration of the CLL-IPI in their cohort.
Overall judgement	No	

CLL-IPI V - Gentile 2016 (Italian cohort)

CLL-IPI V - Gentile 2016 (I Study characteristics	reacian conorcy
General information	Model and type of study
	Gentile 2016 (Italian multicentre cohort)Validation study
	Secondary citations
	• Gentile 2017
	Language of publication
	• English
	Study design
	retrospective cohort
	Follow-up time
	median 69.6 months, range not reported
Participants	Number of included persons in the cohort
	858 persons
	Setting
	Italy, multicentre study
	Recruitment period
	• 1985 - 2015
	Age (in years)
	median 65.5 years (range: not reported)

Sex

• 56.2% male



CLL-IPIV - Gentile 2016 (Italian cohort) (Continued)

Stages of disease

- Rai 0: 58.4%; Rai I: 19.8%; Rai II: 15.2%; Rai III: 1.5%; Rai IV: 5.1%
- Binet A: 79.7%; Binet B: 18.8%; Binet C: 1.5%

Treatment

- 130 received chemotherapy
- 174 received chemoimmunotherapy

Inclusion criteria

 All patients with CLL in one of the five centres that were evaluable for CLL-IPI (i.e. fully available data on age, staging, IgHV, ß2m and del17p/TP53)

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- TP53 was replaced by the proxy del(17p).

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

 overall survival: for OS, the time interval was measured from the day of CLL diagnosis until death from all causes or last follow-up.

Additional outcome(s)

 time-to-first-treatment: For TTFT from the day of CLL diagnosis until the start of therapy or last follow-up

Outcome in model development

overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 858 persons, 174 events
- time-to-first-treatment: 858 persons, 304 events

Which model was used?

· simplified model

Was the model updated?



CLL-IPIV - Gentile 2016 (Italian cohort) (Continued)

n

Performance measures reported?

- Calibration: not reported, survival per risk group and explained variation on outcome available
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- · at 5 years
- percentage of group without event at time point (low risk: 94.5%, intermediate risk: 86.4%, high risk: 74.4%, very high risk: 42.9%)

PROBAST: Applicability

Domain 1: Participant selection

unclear - data was retrospectively collected for a large time period (1985 - 2015). During this time, the
definition of CLL and treatment options have changed. Therefore, the cohort may be very heterogeneous and discrimination for OS may not reflect performance of the model for individuals diagnosed
nowadays.

Domain 2: Predictors

• unclear - predictor TP53 not available and replaced by a proxy, del(17p)

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- · Funding source not reported
- "S.A.P. received funding from Pharmacyclics. All other authors have no conflict of interest to disclose."

Other comments

• identical centres as presented in Gentile 2017

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used: consecutive sampling, but exclusion of individuals without baseline characteristics and FISH analysis from analysis (in total, only 22.5% of individuals were evaluable). This is rated in domain 4.
Domain 2: Predictors	Unclear	Due to the broad time range and multiple participating centres, we doubt that predictor assessments were homogeneous.
Domain 3: Outcome	Yes	Objective standard outcome: ' death, which were abstracted from clinical records at the time of inclusion and updated on an ongoing basis.'
Domain 4: Analysis	No	Consecutive sampling, but exclusion of individuals without baseline characteristics and FISH analysis from analysis (in total, only 22.5% of patients were evaluable). Missing values emerged due to the long time span. Some predictors were not assessed by default at diagnosis until recently. Comparison of baseline sample and included sample similar.
		Calibration not reported, however, survival per risk group available
Overall judgement	No	



CLL-IPI V - Molica 2016 (O-CLL1-GISL)

Study characteristics

General information

Model and type of study

- Molica 2016 (O-CLL1-GISL)
- · Validation study

Secondary citations

• Molica 2017; Molica 2017; Molica 2016

Language of publication

• English

Study design

· prospective cohort

Follow-up time

• median 42 months, 2038 person-years

Participants

Number of included persons in the cohort

• 337 persons

Setting

· Italy, multicentre study

Recruitment period

• 2007 - not reported

Age (in years)

• median 61 years (range: 33 - 70 years)

Sex

• 57.2% male

Stages of disease

- Rai 0: 77.8%; Rai ≥I: 22.2%
- Binet A: 100%; Binet B/C: 0%

Treatment

- 91 received therapy
- · unclear which therapy

Inclusion criteria

 Newly-diagnosed CLL patients from several Italian Institutions who were seen within 12 months of diagnosis were prospectively enrolled into the OCLL1- GISL protocol.

Exclusion criteria

· not reported



CLL-IPIV - Molica 2016 (O-CLL1-GISL) (Continued)

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- TP53 and del17p as composite

Timing of predictor measurement

• at study entry, within 12 months of diagnosis

Outcome(s)

Primary outcome in study

time-to-first-treatment: the primary endpoint, TTFT, was defined as the interval between the date of registration and the date of initiation of first CLL treatment. Patients underwent sequential monitoring, and the frequency of follow-up visits was individualised according to patient risk; this ranged from 3 to 6 months (median 6 months). All physicians who registered patients in this observational database stated that they had used the NCI-WG guidelines as a reference criterion for starting therapy. In particular, the absolute lymphocyte count was not used as the sole indicator for treatment. Active disease, requiring therapy, was defined when at least one of the criteria set out in the NCI sponsored Working Group guidelines was satisfied.

Additional outcome(s)

not applicable

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 337 persons, 91 events

Which model was used?

simplified model

Was the model updated?

- no
- new cut-off score for risk groups tested

Performance measures reported?

- · Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- at 5 years



CLL-IPI V - Molica 2016 (O-CLL1-GISL) (Continued)

percentage of group without event at time point (low risk: 76%, intermediate risk: 45%, high risk: 41%, very high risk: not available)

PROBAST: Applicability

Not applicable; outcome did not match primary outcome of the model.

Notes

Funding & conflict of interest

- "Contract grant sponsor: NIH; Contract grant no.: 1R01CA197120-01."; "Contract grant sponsors: AIRC (the Italian Association for Cancer Research), a non-profit organization to FM. "Special Program Molecular Clinical Oncology 5 per mille" n. 9980, 2010/15 and AIRC "Innovative immunotherapeutic treatments of human cancer" n.16695, 2015/18."
- · "Conflict of interest: Nothing to report"

Other comments

• The same model was validated on the same cohort in other publications.

CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort)

Study characteristics

General information

Model and type of study

- · Munoz-Novas, 2018 (Spanish cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

· English

Study design

· retrospective cohort

Follow-up time

• median 68 months, range not reported

Participants

Number of included persons in the cohort

• 696 persons

<u>Setting</u>

· Spain, multicentre study

Recruitment period

• 1989 - 2013

Age (in years)

• median 65.7 years (range: IQR 55.2 - 73.5)

<u>Sex</u>

• 62.8% male



CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort) (Continued)

Stages of disease

- Rai 0: 58.9%; Rai I-IV: 41.1%
- · Binet A: not reported

Treatment

- 137 received treatment
- · unclear which treatment

Inclusion criteria

• all patients diagnosed with CLL

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); Age (continuous: categorical/binary transformation; cut-off 65 years)
- TP53 not assessed, only del(17p) available

Timing of predictor measurement

· not reported, presumably at diagnosis

Outcome(s)

Primary outcome in study

 overall survival: Overall survival (OS) was calculated from the time of diagnosis to death or last follow-up.

Additional outcome(s)

• time-to-first-treatment: Time to first therapy (TTFT) from the date of diagnosis to first treatment or last follow-up.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

• yes (438 participants)

If yes, how was missing data handled?

· complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 258 persons, 47 events
- time-to-first-treatment: 258 persons, 113 events

Which model was used?

• simplified model

Was the model updated?



CLL-IPI V - Muñoz-Novas 2018 (Spanish cohort) (Continued)

no

Performance measures reported?

- · Calibration: not reported
- Discrimination: reported (c-statistic) and ROC curve (S2)

Creation of risk groups?

- yes, 4 risk groups, (cut points at (0-1);(2-3);(4-6); (7-10))
- at 5 years
- percentage of group without event at time point (low risk: 93.6%, intermediate risk: 87.6%, high risk: 67.8%, very high risk: 28.6%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

• unclear - Predictor TP53 not available and replaced by a proxy, del(17p)

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- · Funding not reported
- "The authors declare that there are no conflicts of interest regarding the publication of this article."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data source used: 'A total of 696 unselected CLL patients newly diagnosed and previously untreated from different institutions of the central region of Spain were included in this study.'
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment in this publication. We assume standardised outcome assessment.
Domain 4: Analysis	No	Insufficient number of events
		Participants with missing values not included in model - consecutive sampling of routine patients, therefore around 2/3 of patients with missing data
		Calibration not reported
Overall judgement	No	



CLL-IPI V - Rani 2018 (Indian cohort)

Study characteristics

General information

Model and type of study

- · Rani 2018 (Indian cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

unclear

Follow-up time

• median 40.5 months (range: 1 - 215 months)

Participants

Number of included persons in the cohort

· 198 persons

Setting

• India, unclear if single or multicentre study

Recruitment period

· not reported

Age (in years)

• median 60 years (range: not reported)

Sex

• 77% male

Stages of disease

- Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.8%; Rai III: 14.1%; Rai IV: 15.2%
- Binet A: not reported

Treatment

- 62 received chlorambucil-based therapy
- 56 received rituximab-based therapy
- 20 received other therapies

Inclusion criteria

· treatment-naive CLL persons according to Hallek 2008

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



CLL-IPIV - Rani 2018 (Indian cohort) (Continued)

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- only TP53 deletions were used to document TP53 aberrations because of unavailability of TP53 mutation status for all the patients.

Timing of predictor measurement

· not reported

Outcome(s)

Primary outcome in study

 overall survival: OS was defined as the time from the date of diagnosis to date of death or date of last follow-up.

Additional outcome(s)

time-to-first-treatment: TTFT was defined as the time from the date of diagnosis to date of commencement of first therapy.

Outcome in model development

overall survival

Missing data

Participants with any missing data?

nc

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- · overall survival: 198 persons, 86 events
- time-to-first-treatment: 140 persons, 89 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups (cut-points at (0-1);(2-3);(4-6); (7-10))
- median survival (low risk: not reached, intermediate risk: 190 years, high risk: 62 years, very high risk: 28 years)

PROBAST: Applicability

<u>Domain 1: Participant selection</u>

• unclear - eligibility criteria were not reported, possibly patients with missing values excluded. We are unsure if the sample is representative for CLL patients.

Domain 2: Predictors



CLL-IPI V - Rani 2018 (Indian cohort) (Continued)

• unclear - TP53 was used as deletion only instead of deletion or mutation.

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "The financial support was provided by the Department of Biotechnology (BT/PR11106/ GBD/27/145/2008, BT/PR15438/MED/30/606/2011 and T/PR8680/AGR/36/754/2013), Ministry of Science and Technology, GOI, and All India Institute of Medical Sciences, New Delhi (8-60/A060/2011/ RS) to RG for carrying out this work."
- "The authors declare that they have no conflict of interest."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Inclusion and exclusion criteria unclear, it seemed like only patients with complete datasets were included. The recruitment period unclear
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way (single-centre study).
Domain 3: Outcome	Unclear	Observation time was short.
Domain 4: Analysis	No	Low number of events; no information on handling of missing values; calibration was not reported.
Overall judgement	No	

CLL-IPI V - Reda 2017 (Milano cohort)

Study characteristics

General information

Model and type of study

- Reda 2017 (Milano cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

English

Study design

· retrospective cohort

Follow-up time



CLL-IPI V - Reda 2017 (Milano cohort) (Continued)

· median 144 months, range not reported

Participants

Number of included persons in the cohort

• 698 persons

Setting

· Italy, single-centre study

Recruitment period

1983 - 2016

Age (in years)

• median 65 years (range: 32 - 91 years)

Sex

• 57% male

Stages of disease

- Rai 0-I: 74%; Rai II-IV: not reported
- Binet A: 82%; Binet B/C: not reported

Treatment

- 190 received Chlorambucil (CHL)
- 108 received fludarabinecyclophosphamide-rituximab (FCR)
- 12 received of atumum abbendamustine (O-Benda)
- 44 received bendamustine-rituximab (BR)
- 65 received alemtuzumab
- 29 received Ibrutinib or Idelalisib

Inclusion criteria

• all patients diagnosed with CLL reclassified by 2008 criteria

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- It was not clear if TP53 was used, or its proxy del(17p) instead.

Timing of predictor measurement

not reported, probably at diagnosis (medical records)

Outcome(s)

Primary outcome in study

• overall survival: OS was defined as time from CLL diagnosis to death.

Additional outcome(s)

• time-to-first-treatment: TTFT was evaluated from the time of diagnosis to first-line therapy start.



CLL-IPI V - Reda 2017 (Mila	no cohort) (Continued) Outcome in model development
	overall survival
Missing data	Participants with any missing data?
	• yes (369 participants)
	If yes, how was missing data handled?
	• unclear
Analysis	Number of participants & number of events (specific time points where reported)
	 overall survival: 329 persons, number of events not reported time-to-first-treatment: 329 persons, number of events not reported
	Which model was used?
	simplified model
	Was the model updated?
	• no
	Performance measures reported?
	 Calibration: not reported Discrimination: not reported
	Creation of risk groups?
	 yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10)) no time point and numerical results reported, see figure 1
PROBAST: Applicability	Domain 1: Participant selection
	• unclear - long observation period (1983 - 2016), no clear eligibility criteria
	Domain 2: Predictors
	 unclear - not much information on predictors reported, mentioning of FISH data but unclear if TP53 or del(17p) was used as predictor
	Domain 3: Outcome
	• low
Notes	Funding & conflict of interest
	 Funding not reported "Authors have no affiliations that they consider to be relevant and important with any organization that to any author's knowledge has a direct interest, particularly a financial interest, in the subject matter discussed."
	Other comments
	Hematology Unit, IRCCS Ca' Granda Ospedale Maggiore Policlinico of Milan, Milano, Italy
Item	Authors' judgement Support for judgement



CLL-IPI V - Reda 2017 (Milano cohort) (Continued)		
Domain 1: Participant se- lection	Unclear	No eligibility criteria reported
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way (single-centre study)
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment in this publication. We assumed standardised outcome assessment
Domain 4: Analysis	No	Unclear how many events and how many patients etc. were included for validation
		Patients with missing values were excluded from analysis.
		No performance measures and survival per group reported (figure 1 not sharp)
Overall judgement	No	

CLL-IPI V - Rigolin 2017 (Ferrera cohort)

Study characteristics			
General information	Model and type of study		
	Rigolin 2017 (Ferrera cohort)Validation study		
	<u>Secondary citations</u>		
	not applicable		
	Language of publication		
	• English		
	Study design		
	retrospective cohort		
	Follow-up time		
	• not reported		
Participants	Number of included persons in the cohort		
	• 335 persons		
	Setting		
	Italy, single-centre study		
	Recruitment period		
	• 2006 - 2016		
	Age (in years)rri		
	• median 68.7 years (range: 33 - 96 years)		
	<u>Sex</u>		



CLL-IPIV - Rigolin 2017 (Ferrera cohort) (Continued)

• 57.9% male

Stages of disease

- · Rai: not reported
- Binet A: 77.9%; Binet B: 14.3%; Binet C: 7.8%

Treatment

• 114 received chemotherapy

Inclusion criteria

· all patients diagnosed with CLL at 'our center'

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- · no changes

Timing of predictor measurement

· not reported

Outcome(s)

Primary outcome in study

• overall survival: OS was calculated from the date of diagnosis until death due to any cause or until the last patient follow-up.

Additional outcome(s)

 time-to-first-treatment: TTFT was calculated as the interval between diagnosis and the start of firstline treatment.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

· unclear or not reported

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• 335 participants, number of events not reported

Which model was used?

· simplified model

Was the model updated?

no



CLL-IPIV - Rigolin 2017 (Ferrera cohort) (Continued)

Performance measures reported?

Calibration: not reportedDiscrimination: not reported

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- no time point and numerical results reported, see figure 1

PROBAST: Applicability

Domain 1: Participant selection

unclear - no eligibility criteria

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "... Fondo di Ateneo per la Ricerca 2013, 2014, 2016 of the University of Ferrara (G.M.R., A.C., M.N.), Programma Ricerca Regione Università 2007-2009 University of Ferrara (G.M.R., A.C.), Programmi di Ricerca di Rilevante Interesse Nazionale (PRIN) 2008 (A.C.), Ricerca Finalizzata (A.C.; project RF-2011-02349712), Ministero ell'Istruzione, dell'Università e della Ricerca PRIN 2015 (A.C.; project 2015ZMRFEA), and AIL Ferrara. E.V. and E.S. are supported by AIL Ferrara."
- "The authors declare no competing financial interests."

Other comments

 The authors provided their primary data and, in addition, a screen capture of the Stata output of their analysis.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	No eligibility criteria
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way (single-centre study)
Domain 3: Outcome	Unclear	Objective standard outcome, however, observation time not reported
Domain 4: Analysis	No	No information on number of events, missing data and missing data handling; no performance measures reported
Overall judgement	No	

CLL-IPI V - Zhu 2018 (Chinese cohort)

Study characteristics

General information Model and type of study



CLL-IPI V - Zhu 2018 (Chinese cohort) (Continued)

- · Zhu 2018 (Chinese cohort)
- Validation study

Secondary citations

· not applicable

Language of publication

Chinese

Study design

· retrospective cohort

Follow-up time

• median 48 months, (range: 1 - 192 months)

Participants

Number of included persons in the cohort

· 215 persons

Setting

· China, single-centre study

Recruitment period

• 2002 - 2017

Age (in years)

• median 60 years (range 16 - 85 years)

Sex

• 66.5% male

Stages of disease

- Rai 0: 16.7%; Rai I-II: 54.0%; Rai III-IV: 29.3%
- Binet A: 44.2%; Binet B: 27.9%; Binet C: 27.9%

Treatment

- 147 received any of the following:
- benzodiazepine
- FCR (fludarabine, cyclophosphamide, rituximab),
- bendamoxetine
- rituximab
- R+FFP+HDMP (rituximab + fresh frozen plasma + high-dose methylprednisolone)

Inclusion criteria

• newly-diagnosed CLL patients treated in our hospital between 2002 and 2017

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



CLL-IPI V - Zhu 2018 (Chinese cohort) (Continued)

- TP53 status (binary; deleted or mutated); IgHV mutational status (continuous: categorical/binary transformation; cut-off not given but usually 98%); ß2-microglobulin concentration (continuous: categorical/binary transformation; binary, cut-off 3.5 mg/L (based on literature)); clinical stage (Rai or Binet) (categorical); age (continuous: categorical/binary transformation; cut-off 65 years)
- · no changes

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

 overall survival: OS time was defined as the interval from a definitive diagnosis to the end point of death or follow-up for any cause.

Additional outcome(s)

time-to-first-treatment: TTFT was defined as the interval from the definitive diagnosis to the initiation
of treatment.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 215 persons, 46 events
- time-to-first-treatment: 215 persons, 147 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: was provided upon request (observed-expected ratio per risk group, calibration plot, Hosmer and Lemeshow test)
- Discrimination: reported (AUC)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-1);(2-3);(4-6); (7-10))
- at 5 years
- percentage of group without event at time point (low risk: 97.6%, intermediate risk: 83.7%, high risk: 67.8%, very high risk: 55.2%)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - no eligibility criteria reported.

Domain 2: Predictors



CLL-IPI V - Zhu 2018 (Chinese cohort) (Continued)

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

· not reported

Other comments

- original study translated by Yuan Chi (Centre for Evidence-Based Chinese Medicine, Beijing University of Chinese Medicine, China)
- the authors provided additional data upon request (calibration plot, O:E ratios per risk group, AUC).

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	No eligibility criteria reported, unclear if patients with missing values were excluded
Domain 2: Predictors	Yes	Well-established predictors used - not explicitly stated, but predictors probably assessed in a similar way
Domain 3: Outcome	Yes	Objective standard outcome
Domain 4: Analysis	No	Number of events low (46)
		No information on missing values
		The authors provided the calibration and discrimination upon request.
Overall judgement	No	

GCLLSG D - Pflug 2014 (GCLLSG)

Study characteristics

General information

Model and type of study

- Pflug 2014 (development cohort)
- Development study

Secondary citations

• Tam 2014

Language of publication

English

Study design

• combined data (e.g. IPD or combination of cohorts); data from 3 RCTs

Follow-up time



GCLLSG D - Pflug 2014 (GCLLSG) (Continued)

• median 63.4 months (range not reported)

Participants

Number of included persons in the cohort

· 2007 persons

Setting

- · multicentre study
- Australia, Austria, Belgium, Czech Republic, Denmark, France, Germany, Israel, Italy, New Zealand, Spain

Recruitment period

1997 - 2006

Age (in years)

• median 60 years (range: 30 - 81 years)

Sex

• 68.4% male

Stages of disease

- Rai 0: 20.8%; Rai I: 18.4%; Rai II: 37.9%; Rai III: 8.6%; Rai IV: 14.3%
- Binet A: 42.6%; Binet B: 37.7%; Binet C: 19.8%

Treatment

- participants were treated with:
- early fludarabine (F)
- F plus cyclophosphamide (FC)
- FC plus rituximab (FCR)

Inclusion criteria

 different inclusion criteria per clinical trial (CLL1, CLL4 and CLL8 trials); common criteria: confirmed diagnosis of B-cell chronic lymphocytic leukaemia, untreated patient, life expectancy above 6 months

Exclusion criteria

• different exclusion criteria per clinical trial (CLL1, CLL4 and CLL8 trials)

Predictors

Number of candidate predictors

- univariable analysis: 20
- multivariable analysis: 18

<u>List of predictors in final model (including cut-points for dichotomised factors)</u>

sex (binary); age (continuous, dichotomisation after univariable analysis); ECOG PS (categorical, dichotomisation after univariable analysis); del(17p) (continuous: categorical/binary transformation); del(11q) (continuous: categorical/binary transformation; literature-based); IgHV mutational status (continuous: categorical/binary transformation; literature-based); s-TK (continuous, dichotomisation after univariable analysis); s-beta2m (continuous, dichotomisation after univariable analysis)

Timing of predictor measurement

· not reported, presumably at study entry

Outcome(s)

Primary outcome in study



GCLLSG D - Pflug 2014 (GCLLSG) (Continued)

overall survival (the main end point of statistical analyses was OS defined as the time between registration/randomisation and death)

Additional outcome(s)

· not applicable

Missing data

Participants with any missing data?

· yes (784 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

· overall survival: 1223 participants, number of events included for analysis unclear

Predictor selection method

• univariable analysis, during multivariable modelling: stepwise regression, (P value or CI)

Statistical method

· Cox proportional hazard model

Simplification of model?

yes

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-2); (3-5); (6-10); (11-14))
- at 5 years
- percentage of group without event at time point (low risk: 95.2 %, intermediate risk: 86.9 %, high risk: 67.6 %, very high risk: 18.7 %)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - the study used RCT data with specific inclusion and exclusion criteria, patient population younger than average

Domain 2: Predictors

• low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- "This manuscript was written on behalf of the German CLL Study Group. Studies CLL1, CLL4, and CLL8 were planned and conducted as investigator-initiated trials by the German CLL Study Group and were supported by research grants from German Cancer Aid, Medac Schering Onkologie, and F. Hoffmann-La Roche. T.D.S. is a clinical scholar of the Leukemia Lymphoma Society."
- "N.P. received Travel Grants from Roche. T.D.S. received research grants from Genentech, Celgene, Glaxo-Smith-Kline, Cephalon, Hospira, and Polyphenon E International. B.E. is a consultant and/or



GCLLSG D - Pflug 2014 (GCLLSG) (Continued)

holds an advisory role for Celgene and Pharmacyclics and has received honoraria and research funding from Roche and Mundipharma. S.S. is a consultant and/or holds an advisory role for Roche and Mundipharma, and received honoraria and research funding from both. H.D. received research grants from Roche. U.J. received honoraria and research funding from Roche. M.H. is a consultant and/or holds an advisory role and received research funding from Roche. J.B., M.A.B., T.E., K.B., G.M., K.G.R., M.J.E., G.H., R.B., A.-M.F., C.-M.W., K.F., and N.E.K. declare no competing financial interests."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used: combination of three RCT cohorts
Domain 2: Predictors	Yes	Not explicitly stated in this publication, but we assumed standard assessment in the context of each RCT
Domain 3: Outcome	Yes	Objective standard outcome, follow-up in the context of each individual RCT
Domain 4: Analysis	No	Univariable selection of predictors
		Bootstrapping, however only for multivariable modeling and not used for correction of optimism; complete case analysis, background sample similar but many missing values; calibration not reported; simplification of the model, thus the assigned weights did not correspond to the results of multivariable modeling.
Overall judgement	No	

GCLLSG V - Molica 2015 (O-CLL1-GISL)

Study characteristics

General information

Model and type of study

- Molica 2015 (O-CLL1-GISL)
- Validation study

Secondary citations

• Molica 2013; Molica 2017; Molica 2017

Language of publication

• English

Study design

· prospective cohort

Follow-up time

• median 42 months (range: 1 - 82 months; 2038 person-years)

Participants

Number of included participants in the cohort



GCLLSG V - Molica 2015 (O-CLL1-GISL) (Continued)

· 337 participants

Setting

· Italy, multicentre study

Recruitment period

• 2007 - not reported

Age (in years)

• median 61 years (range: 33 - 70 years)

Sex

• 57.1% male

Stages of disease

- Rai 0: 77.8%; Rai I-IV: not reported
- Binet A: 100 %

Treatment

- · 91 received treatment
- · unclear which treatment

Inclusion criteria

newly-diagnosed CLL patients from several Italian Institutions who were seen within 12 months of diagnosis were prospectively enrolled into the OCLL1-GISL protocol. The inclusion criteria for CLL diagnosis, employed at the time of study design, were those of the National Cancer Institute (NCI)-sponsored Working Group guidelines (NCI-WG). Patients enrolled did not require therapy according to NCI guidelines (i.e. asymptomatic Binet stage A).

Exclusion criteria

not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- sex (binary); age (continuous, dichotomisation after univariable analysis); ECOG PS (categorical, dichotomisation after univariable analysis); del(17p) (continuous: categorical/binary transformation); del(11q) (continuous: categorical/binary transformation; literature-based); IgHV mutational status (continuous: categorical/binary transformation; literature-based); s-TK (continuous, dichotomisation after univariable analysis); s-beta2m (continuous, dichotomisation after univariable analysis)
- TK was omitted.

Timing of predictor measurement

• at study entry, within 12 months of diagnosis

Outcome(s)

Primary outcome in study

time-to-first-treatment (TTFT was defined as the interval between the date of database registration
and the date of first CLL treatment. Patients underwent sequential monitoring, and the frequency of
follow-up visits was individualised according to patient risk; this ranged from 3 to 6 months (median
6 months). All physicians who registered patients in this observational database stated that they had
used the NCI-WG guidelines as a reference criterion for starting therapy. In particular, the absolute
lymphocyte count was not used as the sole indicator for treatment. Active disease, requiring therapy,
was defined when at least one of the criteria set out in the NCI-sponsored Working Group guidelines
was satisfied).



GCLLSG V - Molica 2015	(O-CLL1-GISL)	(Continued)
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Additional outcome(s)

· not applicable

Outcome in model development

overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 337 participants, 91 events

Which model was used?

· simplified model

Was the model updated?

nc

Performance measures reported?

· Calibration: not reported

· Discrimination: reported (c-statistic)

Creation of risk groups?

no

PROBAST: Applicability

Not applicable; outcome did not match primary outcome of the model.

Notes

Funding & conflict of interest

- "Contract grant sponsor: NIH; Contract grant number: 1R01CA197120-01."; "Contract grant sponsors:
 AIRC (the Italian Association for Cancer Research), a non-profit organization to FM. "Special Program
 Molecular Clinical Oncology 5 per mille" n. 9980, 2010/15 and AIRC "Innovative immunotherapeutic
 treatments of human cancer" n.16695, 2015/18." (Molica 2016)
- "Conflict of interests: nothing to report." (Molica 2016)

Other comments

- The validation of this model in the O-CLL1-GISL cohort appeared in several publications. The c-statistic was reported differently in each of them.
- The authors did not reply to our request for additional information.

GCLLSG V - Pflug 2014 (Mayo cohort)

Study characteristics

General information

Model and type of study

- Pflug 2014 (Mayo clinic cohort, undefined time range)
- Validation study



GCLLSG V - Pflug 2014 (Mayo cohort) (Continued)

Secondary citations

· not applicable

Language of publication

• English

Study design

· prospective cohort

Follow-up time

· median 57 months (range not reported)

Participants

Number of included participants in the cohort

· 676 participants

Setting

· USA, multicentre study

Recruitment period

· not reported

Age (in years)

• median 61.5 years (range: 32 - 89 years)

Sex

• 67% male

Stages of disease

- Rai 0: 57.1%; Rai I: 34%; Rai II: 5.9%; Rai III: 1%; Rai VI: 1.9%
- not reported

Treatment

- · 190 received treatment
- · unclear which treatment

Inclusion criteria

• consecutive series of 676 newly-diagnosed, prospectively-followed CLL patients cared for at Mayo Clinic who had baseline data on all considered variables except s-TK and/or s-B2M available and who had stored serum collected less than 36 months (median, 1 month) of diagnosis available for s-TK and s-B2M analysis.

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

sex (binary); age (continuous, dichotomisation after univariable analysis); ECOG PS (categorical, dichotomisation after univariable analysis); del(17p) (continuous: categorical/binary transformation); del(11q) (continuous: categorical/binary transformation; literature-based); IgHV mutational status (continuous: categorical/binary transformation; literature-based); s-TK (continuous, dichotomisation



GCLLSG V - Pflug 2014 (Mayo cohort) (Continued)

after univariable analysis); serum ß2-microglobulin (continuous, dichotomisation after univariable analysis)

• TK from frozen blood samples within 36 months of diagnosis; different assessment methods for beta2m than in development study

Timing of predictor measurement

· at study entry

Outcome(s)

Primary outcome in study

overall survival: the main end point of statistical analyses was OS defined as the time between registration/randomisation and death.

Additional outcome(s)

• time-to-first-treatment: TFS was calculated from the date of diagnosis to the start of the first CLL treatment.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

· yes (3 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 673 participants, 85 events
- time-to-first-treatment: 673 participants, 190 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-2); (3-5); (6-10); (11-14)) ·
- at 5 years
- percentage of group without event at time point (low risk: 95.2%, intermediate risk: 91.4%, high risk: 71.7%, very high risk: 13.6%)

PROBAST: Applicability

<u>Domain 1: Participant selection</u>

• unclear - unclear recruitment period, data availability as inclusion criterion

Domain 2: Predictors

• low - differences in the measurement of one predictor, which is usually not available in the USA as compared to Germany (serum thymidine kinase): interassay calibration



GCLLSG V - Pflug 2014 (Mayo cohort) (Continued)

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "This manuscript was written on behalf of the German CLL Study Group. Studies CLL1, CLL4, and CLL8 were planned and conducted as investigator-initiated trials by the German CLL Study Group and were supported by research grants from German Cancer Aid, Medac Schering Onkologie, and F. Hoffmann-La Roche. T.D.S. is a clinical scholar of the Leukemia Lymphoma Society."
- "N.P. received Travel Grants from Roche. T.D.S. received research grants from Genentech, Celgene, Glaxo-Smith-Kline, Cephalon, Hospira, and Polyphenon E International. B.E. is a consultant and/or holds an advisory role for Celgene and Pharmacyclics and has received honoraria and research funding from Roche and Mundipharma. S.S. is a consultant and/or holds an advisory role for Roche and-Mundipharma, and received honoraria and research funding from both. H.D. received research grants from Roche. U.J. received honoraria and research funding from Roche. M.H. is a consultant and/or holds an advisory role and received research funding from Roche. J.B., M.A.B., T.E., K.B., G.M., K.G.R., M.J.E., G.H., R.B., A.-M.F., C.-M.W., K.F., and N.E.K. declare no competing financial interests."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Only patients with available data included: " who had baseline data on all considered variables except s-TK and/or s-b2m available and who had stored serum collected > 36 months"
Domain 2: Predictors	Yes	Standard laboratory measures used. Where s-TK and s-B2M were not available, blood samples were shipped to Germany and analysed.
Domain 3: Outcome	Yes	Objective standard outcome - no clear description of outcome assessment - we assumed that assessment of the objective outcome OS was similar and therefore did not rate as high risk (no info on e.g. registry, frequency of follow-up, etc.)
Domain 4: Analysis	No	Low number of events; patients with missing data were not enrolled in the study, rated in domain 1; calibration not reported
Overall judgement	No	

GCLLSG V - Rani 2018 (Indian cohort)

Study characteristics

General information

Model and type of study

- · Rani 2018 (Indian cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication



GCLLSG V - Rani 2018 (Indian cohort) (Continued)

English

Study design

unclear

Follow-up time

• median 40.5 months (range: 1 - 215 months)

Participants

Number of included participants in the cohort

• 198 participants

Setting

· India, unclear if single or multicentre study

Recruitment period

not reported

Age (in years)

• median 60 years (range not reported)

Sex

• 77% male

Stages of disease

- Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%
- Binet A/B/C: not reported

<u>Treatment</u>

- 62 received chlorambucil-based therapy
- 56 received rituximab-based therapy
- · 20 received other therapies

Inclusion criteria

• treatment-naive CLL patients according to Hallek 2008

Exclusion criteria

· not reported

Predictors

 $\underline{\text{List of predictors used for validation (and changes between original predictors and predictors in validation study)}$

sex (binary); age (continuous, dichotomisation after univariable analysis); ECOG PS (categorical, dichotomisation after univariable analysis); del(17p) (continuous: categorical/binary transformation); del(11q) (continuous: categorical/binary transformation; literature-based); IgHV mutational status (continuous: categorical/binary transformation; literature-based); s-TK (continuous, dichotomisation after univariable analysis); s-beta2m (continuous, dichotomisation after univariable analysis)

Timing of predictor measurement

· not reported

Outcome(s)

Primary outcome in study



GCLLSG V - Rani 2018 (Indian cohort) (Continued)

time-to-first-treatment (TTFT was defined as the time from the date of diagnosis to date of commencement of first therapy)

Additional outcome(s)

· not applicable

Outcome in model development

overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 140 participants, 89 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at (0-2); (3-5); (6-10); (11-14)
- median survival (low risk: not reached, intermediate risk: 36 years, high risk: 4 years, very high risk: 0.01 years)

PROBAST: Applicability

Not applicable, outcome did not match primary outcome of the model

Notes

Funding & conflict of interest

- "The financial support was provided by the Department of Biotechnology (BT/PR11106/GBD/27/145/2008, BT/PR15438/MED/30/606/2011 and T/PR8680/AGR/36/754/2013), Ministry of Science and Technology, GOI, and All India Institute of Medical Sciences, New Delhi (8-60/A060/2011/RS) to RG for carrying out this work."
- "The authors declare that they have no conflict of interest."

Other comments

 Graphs were provided to obtain survival at a specific time point, however, these graphs were printed very small.

GIMEMA D - Molica 2005 (GIMEMA cohort)

Study characteristics



GIMEMA D - Molica 2005 (GIMEMA cohort) (Continued)

General information

Model and type of study

- Molica 2005 (GIMEMA cohort)
- · Development study

Secondary citations

· not applicable

Language of publication

· English

Study design

· retrospective cohort

Follow-up time

• median 54 months, range 4 - 309 months

Participants

Number of included participants in the cohort

• 1138 participants

Setting

· Italy, multicentre study

Recruitment period

• 1991 - 2000

Age (in years)

• median 65 years (range: 27 - 100 years)

Sex

• 54% male

Stages of disease

- Rai 0: 77.2 %; Rai I-III: 22.7%
- Binet A: 100%

Treatment

- 133 received chlorambucil
- 20 received fludarabine alone or in association with cyclophosphamide
- 7 received CHOP-like regimens
- 40 received other therapies (CVP, cyclophosphamide with or without prednisone)

Inclusion criteria

 All previously untreated B-cell CLL patients in Binet stage A whose diagnosis was immunologically confirmed (CD5 +/ Smlg weak) and who were observed at different GIMEMA centres during the period 1991 – 2000 were considered eligible for this study.

Exclusion criteria

• inadequate follow-up

Predictors

Number of candidate predictors



GIMEMA D - Molica 2005 (GIMEMA cohort) (Continued)

- univariable analysis: 10 or more
- multivariable analysis: 6

<u>List of predictors in final model (including cut-points for dichotomised factors)</u>

 Lymphocyte doubling time (LDT, continuous: categorical/binary transformation; dichotomised to above and below 12 months); absolute peripheral blood lymphocytosis (continuous: categorical/binary transformation; two categories, above and below 30x10^9/l); Rai stage (categorical; 0 vs. I-III); gender (binary)

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

progression-free survival (progressive disease was defined on the basis of a shift to a more advanced clinical stage (i.e. from A to B or C) and/or LDT larger than 12 months. For those individuals who did not change Binet stage nor doubled their lymphocyte count, disease progression was considered when one of the following conditions was met: decrease in haemoglobin larger than 3 g/dL, splenomegaly larger than 6 cm below the left costal margin, massive lymphadenopathy (i.e. larger than 8 cm) or progressive lymphadenopathy and/or peripheral blood lymphocytosis more than 100x10^9 /l)

Additional outcome(s)

not applicable

Missing data

Participants with any missing data?

• yes (545 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

 progression-free survival: 593 participants, 235 events in overall sample, unclear number of events in sample used for modeling

Predictor selection method

• univariable analysis, during multivariable modelling: unclear if stepwise regression was used (criteria were P value or CI)

Statistical method

• Cox proportional hazard model, stratification of the resulting score by gender

Simplification of model?

yes

Performance measures reported?

- Calibration: not reported
- · Discrimination: not reported

Creation of risk groups?

- yes, 3 risk groups (risk groups were: no risk factors; one risk factor; more than one risk factor)
- at 10 years



GIMEMA D - Molica 2005 (GIMEMA cohort) (Continued)

• percentage of group without event at time point (low risk: 67.8%, intermediate risk: 41%, high risk: 24.8%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

not reported

Other comments

• The model development characteristics were not well reported.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used (retrospective cohort); 1.7% of patients were excluded due to inadequate follow-up. We rated this as low proportion of exclusions.
Domain 2: Predictors	Yes	The predictors used seem to be routine demographic and laboratory information, and although there was no explicit information, we think that the methods were consistent.
Domain 3: Outcome	No	Knowledge of the predictor was used both in the definition of progression and the model (peripheral blood lymphocytes).
Domain 4: Analysis	No	Individuals with missing data were excluded: "The outcome of the 593 patients with complete data was different from the outcome of the 545 patients with incomplete data (10 year PFS 55.5% vs. 70.2%). The reasons for this difference rely on the higher number of patients in Rai stages I – III ($P = 0.01$) and increased serum levels of b2-m ($P = 0.003$) found in the subgroup with complete data"; univariable selection of predictors; no weighting of predictors: "These differences in the RR did not prevent us from constructing a risk score simply by adding the negative factors present in a single patient at the time of diagnosis."; no model performance measures reported
Overall judgement	No	

GIMEMA V - González Rodríguez 2009 (Cabueñes coh.)

Study characteristics

General information

Model and type of study

- Gonzalez Rodriguez 2009 (Hospital de Cabuenes)
- Validation study



GIMEMA V - González Rodríguez 2009 (Cabueñes coh.) (Continued)

Secondary citations

· not applicable

Language of publication

• Spanish

Study design

· retrospective cohort

Follow-up time

· not reported

Participants

Number of included participants in the cohort

· 265 participants

Setting

· Spain, single-centre study

Recruitment period

1997 - 2007

Age (in years)

• median 71.7 years (range: 42 - 94 years)

Sex

• 58.5% male

Stages of disease

- modified Rai low: 61.2%; Rai intermediate: 29.3%; Rai high: 9.5%
- Binet A: 76.8%; Binet B: 14.8%; Binet C: 8.4%

Treatment

· not reported

Inclusion criteria

 Only cases with an increase of more than 5*10^9/l lymphocytes with morphology and immunophenotype of B-CLL and with Royal Mardsen Scoring System greater than 3, and cases of CLL confirmed by lymph node biopsy have been included. The inclusion criteria were based on the initial data at the time of diagnosis.

Exclusion criteria

not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- Lymphocyte doubling time (LDT, continuous: categorical/binary transformation; dichotomised to above and below 12 months); absolute peripheral blood lymphocytosis (continuous: categorical/binary transformation; two categories, above and below 30x10^9/l); Rai stage (categorical; 0 vs. I-III); gender (binary)
- · no changes

Timing of predictor measurement



GIMEMA V - González Rodríguez 2009 (Cabueñes coh.) (Continued)

· at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: to estimate the overall survival (OS), the date of diagnosis and the date of death for any cause or the final date of the study were considered.

Additional outcome(s)

• time-to-first-treatment: treatment-free survival was considered as the time elapsed from diagnosis until the start date of the treatment.

Outcome in model development

· progression-free survival

Missing data

Participants with any missing data?

• yes (8 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 257 participants, 76 events
- time-to-first-treatment: 257 participants, 67 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (AUC)

Creation of risk groups?

- · yes, 3 risk groups
- at 5 years
- percentage of group surviving at 5 years (low risk: 79%, intermediate risk: 67%, high risk: 79%)

PROBAST: Applicability

Domain 1: Participant selection

low

<u>Domain 2: Predictors</u>

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

• This work was partially funded through research project FIS06/ 0841 from the Health Research Fund.



GIMEMA V - González Rodríguez 2009 (Cabueñes coh.) (Continued)

Other comments

- The original study was translated by Leonardo Perales-Guerrero (medical student, Universidad de Guadalajara, Mexico).
- The authors could not be contacted since the email address of the contact author as stated in the publication was no longer valid.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used: retrospective cohort
Domain 2: Predictors	Yes	The predictors used seem to be routine demographic and laboratory information, and although there was no explicit information, we think that the methods are consistent.
Domain 3: Outcome	Unclear	No median observation time; standard outcome, but treatment indication not described
Domain 4: Analysis	No	Only small amount of missing outcome data (n = 8); discrimination reported in form of AUC, however, time point not clear (probably 5 years); calibration not reported
Overall judgement	No	

MDACC 2007 D - Wierda 2007 (MDACC)

Study characteristics			
General information	Model and type of study		
	 Wierda 2007 (development, MDACC 1981-2004) Development study 		
	<u>Secondary citations</u>		
	not applicable		
	Language of publication		
	• English		
	Study design		
	retrospective cohort		
	Follow-up time		
	 median 58.8 months, 95% CI: 55.2 - 61.2 		
Participants	Number of included participants in the cohort		
	• 1674 participants		
	Setting		

• US (Texas), single-centre study



MDACC 2007 D - Wierda 2007 (MDACC) (Continued)

Recruitment period

• 1981 - 2004

Age (in years)

• median 58 years (range: 0 - 90 years)

Sex

• 1029 male participants

Stages of disease

- Rai 0: 469 individuals; Rai I: 739 individuals; Rai II: 235 individuals; Rai III: 93 individuals; Rai IV: 127 individuals
- Binet A: 1019 individuals; Binet B: 470 individuals; Binet C: 168 individuals

Treatment

- 902 received treatment
- · unclear which treatment

Inclusion criteria

 Previously untreated patients who presented for initial evaluation to MDACC from August 1981 through August 2004 were included in this analysis.

Exclusion criteria

· not reported

Predictors

Number of candidate predictors

- univariable analysis: 18
- multivariable analysis: 16

<u>List of predictors in final model (including cut-points for dichotomised factors)</u>

Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); sex (binary)

Timing of predictor measurement

at study entry

Outcome(s)

Primary outcome in study

• overall survival (the time interval was measured from the day of presentation to MDACC until death or last follow-up. Death from all causes was included).

Additional outcome(s)

· not applicable

Missing data

Participants with any missing data?

• yes (57 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)



MDACC 2007 D - Wierda 2007 (MDACC) (Continued)

• overall survival: 1617 participants, 437 events

Predictor selection method

• univariable analysis, during multivariable modelling: backward elimination (P value or CI)

Statistical method

• Cox proportional hazard model

Simplification of model?

yes

Performance measures reported?

- · Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3);(4-7);(above 8))
- at 5 years
- percentage of group without event at time point (low risk: 0.97%, intermediate risk: 0.8%, high risk: 0.55%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- · Funding not reported
- "The authors declare no competing financial interests."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Retrospective cohort, consecutive sampling
Domain 2: Predictors	Yes	Routine clinical data used: although the recruitment period was very broad, standard clinical data (gender, age, Rai stage, ALC and beta-2 microglobulin) have been included for modelling
Domain 3: Outcome	Yes	Objective standard outcome: no clear description of outcome assessment; we assumed that assessment of the objective outcome OS was similar and therefore did not rate as high risk.



MDACC 2007 D - Wierda 2007 (MDACC) (Continued)

Domain 4: Analysis

Ye

Number of missing values and therefore the number of patients excluded from analysis was low; continuous factors handled appropriately; performance measures of original model reported; bootstrap correction applied; original formula and a simplified score provided; univariable selection of factors, however, nearly all factors were included in multivariable modelling as well.

Overall judgement

Yes

MDACC 2007 V - Bulian 2011 (Italian-Swiss)

Study characteristics

General information

Model and type of study

- Bulian 2011 (Italian-Swiss cohort)
- · Validation study

Secondary citations

not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

• median 63.6 months, range not reported

Participants

Number of included participants in the cohort

• 1480 participants

Setting

• Italy, Switzerland, multicentre study

Recruitment period

• 1996 - 2011

Age (in years)

• median 65 years (range: 21 - 94 years)

Sex

• 58.2% male

Stages of disease

- Rai 0-II: not reported; Rai III/IV: 7.2%
- Binet A: 74.6%; Binet B: 18.4%; Binet C: 6.9%

Treatment

• 306 received chemotherapy



MDACC 2007 V - Bulian 2011 (Italian-Swiss) (Continued)

- · 129 received Chemoimmunotherapy
- · 42 received FCR
- 49 received FR
- 38 received R-other
- · 40 with missing data

Inclusion criteria

 'The series included all cases observed at each centre over a given time period, although each centre allegedly failed to observe all the incident CLL cases.'

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)
- · no changes

Timing of predictor measurement

 not explicitly reported, but probably at diagnosis, as this was a retrospective cohort using standard demographic and clinical information

Outcome(s)

Primary outcome in study

· overall survival

Additional outcome(s)

• time-to-treatment ('Patient treatment was started according to NCI-WG indications.')

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• overall survival: 1037 participants, 151 events

Which model was used?

• The authors tried to use the nomogram, but had problems due to the low quality of the graph in the development publication; additionally, they used the index score.

Was the model updated?

- ves
- 'An additional model was developed by replacing the dichotomous variables Rai staging (0–I–II vs. III–IV) with the three-level Binet staging.'

Performance measures reported?



MDACC 2007 V - Bulian 2011 (Italian-Swiss) (Continued)

Calibration: not reportedDiscrimination: not reported

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3), (4-7), (>7))
- · at 5 years
- percentage of group without event at time point (low risk: 97%, intermediate risk: 89%, high risk: 53%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "Ministero della Salute (Ricerca Finalizzata I.R.C.C.S. and 'Alleanza Contro il Cancro'), Rome; Associazione Italiana contro le Leucemie, linfomi e mielomi (A.I.L.), Venezia Section, Pramaggiore Group; Ricerca Scientifica Applicata, Regione Friuli Venezia Giulia, Trieste ('Linfonet' project); Associazione Italiana per la Ricerca sul Cancro (Investigator Grant IG-8701), Milan, Italy; Helmut Horten Foundation, San Salvatore Foundation, Fondazione per la Ricerca e la Cura sui Linfomi (Lugano, Switzerland); Swiss Cancer League (Krebsliga Schweiz) grant KLS-01690-03-2005."
- "The authors declare no competing financial interests."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	"Clinical and biological data of 1480 untreated CLL patients were retrospectively collected from 8 centres. The series included all cases observed at each centre over a given time period,"
Domain 2: Predictors	Yes	Measurements were taken in 8 different centres. However, the authors standardised the predictors so this should not be of influence.
Domain 3: Outcome	Yes	Objective standard outcome: no definition of outcome and assessment - we assumed that definition and assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	Exclusion of individuals with missing values: "Some CLL cases with missing values were excluded from multivariate models. The death rate in this group was higher. The lack of inclusion of a number of individuals with possible worse prognosis could partly justify the observed bias in nomogram prediction."; expected OS estimated, but only point score: no discrimination calculated, because estimation based on nomogram not possible (quality of graph)
Overall judgement	No	



MDACC 2007 V - Gentile 2014 (Italian cohort)

Study characteristics

General information

Model and type of study

- Gentile 2014 (Italian multicentre cohort)
- Validation study

Secondary citations

• Gentile 2016

Language of publication

· English

Study design

· retrospective cohort

Follow-up time

• median 68.4 months, range not reported

Participants

Number of included participants in the cohort

· 1502 indviduals

Setting

· Italy, multicentre study

Recruitment period

• 1983 - 2013

Age (in years)

• median 67 years (range: not reported)

Sex

• 55.7% male

Stages of disease

- Rai 0: 56.5%; Rai I: 21.4%; Rai II: 15%; Rai III: 2.6%; Rai IV: 4.5%
- Binet A: 82.9%; Binet B: 11.6%; Binet C: 5.5%

Treatment

- · 337 received chemotherapy
- 142 received chemoimmunotherapy

Inclusion criteria

• The CLL databases of four Italian centres, which included all patients diagnosed with CLL since 1983, were utilised for research purposes.

Exclusion criteria

Missing data, ß2m

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



MDACC 2007 V - Gentile 2014 (Italian cohort) (Continued)

Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary);
 Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)

Timing of predictor measurement

· at study entry

Outcome(s)

Primary outcome in study

• overall survival: for OS, the time interval was measured from the day of CLL diagnosis until death from all causes or last follow-up.

Additional outcome(s)

 time-to-first-treatment: for TFS, it was from the day of CLL diagnosis until therapy start or last follow-up.

Outcome in model development

overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 1502 participants, 277 events
- time-to-first-treatment: 1502 participants, 479 events

Which model was used?

· Authors used both nomogram and the index score, and also compared with previous validations.

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3);(4-7);(above 7))
- at 5 years
- percentage of group without event at time point (low risk: 89%, intermediate risk: 82%, high risk: 42%)

PROBAST: Applicability

Domain 1: Participant selection

unclear - very long period of retrospective enrolment: patients were probably treated differently. Inclusion of cMBL patients previously classified as CLL - sensitivity analysis showed no difference in discrimination.

<u>Domain 2: Predictors</u>

low



MDACC 2007 V - Gentile 2014 (Italian cohort) (Continued)

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- Funding not reported
- "The authors indicated no potential conflicts of interest."

Other comments

• 4 of the 5 Italian centres overlapped with the cohort in the secondary citation. This paper was chosen as primary because more information was provided.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Retrospective cohort: " which included all patients diagnosed with CLL since 1983, were utilized for research purposes."
Domain 2: Predictors	Yes	Routine clinical data used: although the recruitment period was very broad, standard clinical data (gender, age, Rai stage, ALC and beta-2 microglobulin) have been included for modelling.
Domain 3: Outcome	Yes	Objective standard outcome: no definition of outcome and assessment - we assumed that definition and assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	Individuals with missing values were excluded. Although background characteristics were similar, we rated this as high risk because nearly half of all patients were excluded from analysis; measure for calibration not reported, however, reporting of survival per risk group
Overall judgement	No	

MDACC 2007 V - Gentile 2016 (Mayo cohort)

Study characteristics

General information

Model and type of study

- Gentile 2016 (Mayo clinic cohort)
- · Validation study

Secondary citations

• Gentile 2016; Shanafelt 2009

Language of publication

• English

Study design

· prospective cohort

Follow-up time



MDACC 2007 V - Gentile 2016 (Mayo cohort) (Continued)

· median 86.4 months, range not reported

Participants

Number of included participants in the cohort

• 506 participants

Setting

· USA, single-centre study

Recruitment period

• 2001 - 2008 (correspondence with authors)

Age (in years)

• median 62.5 years (range: 36 - 89 years)

Sex

• 68.6% male

Stages of disease

- Rai 0: 62.5%; Rai I: 28.9%; Rai II: 5.1%; Rai III: 1.2%; Rai IV: 2.4%
- · Binet A: not reported

Treatment

- 213 received treatment
- · unclear which treatment

Inclusion criteria

not reported

Exclusion criteria

not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary);
 Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)

Timing of predictor measurement

at diagnosis

Outcome(s)

Primary outcome in study

 overall survival: for OS, the time interval was measured from the day of CLL diagnosis until death from all causes or last follow-up.

Additional outcome(s)

• time-to-first-treatment: for TTFT from the day of CLL diagnosis until the start of therapy or last follow-up

Outcome in model development

· overall survival



MDACC 2007 V - Gentile 2016 (Mayo cohort) (Continued)

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Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 506 participants, 114 events
- time-to-first-treatment: 506 participants, 213 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: reported
- · Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3);(4-7);(above 7))
- · at 5 years
- percentage of group without event at time point (low risk: 98.3%, intermediate risk: 84.4%, high risk: 15.6%)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - no clear eligibility criteria

Domain 2: Predictors

• low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- · Funding not reported
- "S.A.P. received funding from Pharmacyclics. All other authors have no conflict of interest to disclose."

Other comments

• The corresponding author provided additional information to describe the cohort (recruitment period, overlap of cohort with other publications)

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Missing values as exclusion reason: 'The validation data set consisted of a consecutive series of 506 newly-diagnosed CLL patients, prospectively followed



MDACC 2007 V - Gentile 20	D16 (Mayo cohort) (Continued)	at the Mayo Clinic, with baseline data available for all variables considered for both CLL-IPI and the MDACC score.'
Domain 2: Predictors	Yes	Routine clinical data used, prospective follow-up, relatively recent cohort (2001-2008)
Domain 3: Outcome	Yes	Objective standard outcome, clear definition and assessment: 'death, which were abstracted from clinical records at the time of inclusion and updated on an ongoing basis.'
Domain 4: Analysis	Unclear	Individuals with missing data were excluded from the study, rated in domain 1. Calibration not reported; survival per risk groups was available.
Overall judgement	No	

MDACC 2007 V - González Rodríguez (Cabueñes)

a			
Stud	v cha	ıracte	ristics

General information

Model and type of study

- Gonzalez Rodriguez 2009 (Cabuenes cohort)
- Validation study

Secondary citations

• not applicable

Language of publication

Spanish

Study design

· retrospective cohort

Follow-up time

· not reported

Participants

Number of included participants in the cohort

· 265 participants

Setting

• Spain, single-centre study

Recruitment period

• 1997 - 2007

Age (in years)

• median 71.7 years (range: 42 - 94 years)

Sex

· not reported



MDACC 2007 V - González Rodríguez (Cabueñes) (Continued)

Stages of disease

- reported as Rai modified stages; low risk: 61.2%; intermediate: 29.3%; high risk: 9.5%
- Binet A: 76.8%; Binet B: 14.8%; Binet C: 8.4%

Treatment

- 67 received treatment
- · unclear which treatment

Inclusion criteria

• Only cases with an increase of more than $5 \times 10^9/l$ lymphocytes with morphology and immunophenotype of B-CLL and with Royal Mardsen Scoring System greater than 3, and cases of CLL confirmed by lymph node biopsy have been included. The inclusion criteria were based on the initial data at the time of diagnosis.

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)
- · no changes

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: to estimate overall survival, the diagnosis date was considered and the date of death by any cause or the final date of the study

Additional outcome(s)

• time-to-first-treatment: treatment-free survival was considered as the time elapsed from diagnosis until the date of treatment initiation.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

• yes (8 participants)

If yes, how was missing data handled?

complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 257 participants, 76 events
- time-to-first-treatment: 257 participants, 67 events

Which model was used?

• index score (simplified model)



MDACC 2007 V - González Rodríguez (Cabueñes) (Continued)

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (AUC)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3);(4-7);(above 7))
- at 5 years
- percentage of group without event at time point (low risk: 87%, intermediate risk: 75%, high risk: 29%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- This work was partially funded through research project FIS06/0841 from the Health Research Fund.
- · Conflict of interest not reported

Other comments

- original study translated by Leonardo Perales-Guerrero (Medical student, Universidad de Guadalajara, Mexico)
- · data extraction from both the original and translated study

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Retrospective cohort, consecutive sampling: 'Un total de 265 pacientes se diagnosticaron de LLCB en el periodo de 10 anos.'
Domain 2: Predictors	Yes	Routine clinical data assessed at diagnosis
Domain 3: Outcome	Yes	Objective standard outcome: no definition of outcome and assessment - we assumed that definition and assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	Low number of events; individuals with missing values excluded from model- ling, however this number was small; calibration not reported
Overall judgement	No	



MDACC 2007 V - Molica 2010 (GIMEMA cohort)

Study characteristics

General information

Model and type of study

- Molica 2010 (GIMEMA cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

• median 41.5 months, range not reported

Participants

Number of included participants in the cohort

- 1158 participants
- characteristics only reported for 310 participants with complete data

Setting

• Italy, multicentre study

Recruitment period

• 1991 - 2000

Age (in years)

• median 64 years (range: not reported)

Sex

• 56.1% male

Stages of disease

- Rai 0: 67%; Rai I-III: 32.9%; Rai III/IV: 0%
- Binet A: 100%

Treatment

- 51 received therapy
- · unclear which treatment

Inclusion criteria

previously untreated CLL participants in Binet stage A whose diagnosis was immunologically confirmed (CD5+/Smlg weak) and who were observed at different GIMEMA primary haematology centres during the period 1991–2000

Exclusion criteria

not reported



MDACC 2007 V - Molica 2010 (GIMEMA cohort) (Continued)

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); sex (binary)
- · cut-off for Rai was adapted in the score

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• time-to-first-treatment: active, therapy-requiring disease was defined by very detailed criteria.

Additional outcome(s)

· overall survival: definition not reported

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 310 participants, number of events not reported
- time-to-first-treatment: 310 participants, 51 events

Which model was used?

• Authors used both nomogram and index score, and also compared with previous validations.

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- · Discrimination: not reported

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3);(4-7);(above 7))
- at 5 years
- percentage of group without event at time point (low risk: 99.3%, intermediate risk: 88.2%, high risk: no individuals in this category)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors



MDACC 2007 V - Molica 2010 (GIMEMA cohort) (Continued)

high - the authors changed the cut-points for the predictor 'Rai stage'. Instead of a binary variable stage
 0-II vs II-IV, Molica 2010 used Rai 0 vs. Rai I-II

Domain 3: Outcome

• High - outcome did not match primary outcome of the model. Survival per risk group reported

Notes

Funding & conflict of interest

- · Funding not reported
- · "The authors reported no potential conflicts of interest."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Retrospective cohort, inclusion of complete database: "The GIMEMA CLL database includes information on previously untreated CLL patients in Binet stage A whose diagnosis was immunologically confirmed".
Domain 2: Predictors	Yes	Although individuals came from 25 different centres, predictors were routine clinical data.
Domain 3: Outcome	Unclear	The median observation time was below 5 years (41.5 months); outcome time-to-first-treatment was not the primary outcome of the validated model.
Domain 4: Analysis	No	Individuals with missing values were excluded from the analysis ("The outcome of the 593 patients with complete data was different from the outcome of the 545 patients with incomplete data"); no performance measures reported
Overall judgement	No	

MDACC 2007 V - Molica 2015 (O-CLL1-GISL)

Study characteristics

General information

Model and type of study

- Molica 2015 (O-CLL1-GISL)
- Validation study

Secondary citations

 Gentile 2016; Molica 2010; Molica 2017 (leukemia & lymphoma); Molica 2017 (American Journal of Hematology)

Language of publication

English

Study design

· prospective cohort



MDACC 2007 V - Molica 2015 (O-CLL1-GISL) (Continued)

Follow-up time

· median 42 months, range not reported

Participants

Number of included participants in the cohort

· 337 participants

Setting

· Italy, multicentre study

Recruitment period

• 2007 - not reported

Age (in years)

• median 61 years (range: 33 - 70 years)

Sex

• 57.1% male

Stages of disease

- Rai 0: 77.5%; Rai I-II: 22.5%
- Binet A: 100%; Binet B/C: not applicable

Treatment

- · 91 received treatment
- · unclear which treatment

Inclusion criteria

• Established diagnosis of B-CLL by NCI criteria, performed by local haematologist (diagnosis will be confirmed by the biological review committee according to flow cytometry analysis (positive clusters of differentiation antigen 5 (CD5), 19 (CD19),23 (CD23)). Age above 18 years and below 70 years. Eastern Cooperative Oncology Group (ECOG) below 2. Binet stage A. Diagnosis performed within 12 months before inclusion in the study. Patients who do not necessitate therapy by NCI guidelines (watch-andwait policy). Shipment of peripheral blood sample to centralised laboratory for biological assessment. Clinical data including baseline information on disease localisation and laboratory parameters at staging and assurance of follow-up updating for at least 3 years are requested. Written informed consent.

Exclusion criteria

Patients with CLL whose diagnosis exceeded 12 months before registration. Patients with leukaemic
phase of lymphoproliferative disorders of B origin CD5- and/or CD23- according to flow cytometry
analysis. Clinical Binet stage B or C. Patients who necessitated therapy according to NCI guidelines
(no watch-and-wait policy). Age above 70 years. Without a written informed consent.

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)
- · no changes

Timing of predictor measurement

• at study entry, within 12 months of diagnosis



MDACC 2007 V - Molica 2015 (O-CLL1-GISL) (Continued)

Outcome(s)

Primary outcome in study

• time-to-first-treatment: the primary endpoint, TTFT, was defined as the interval between the date of database registration and the date of first CLL treatment (individuals underwent sequential monitoring, and the frequency of follow-up visits was individualised according to patient risk; this ranged from 3 to 6 months (median 6 months). All physicians who registered patients in this observational database stated that they had used the NCI-WG guidelines as a reference criterion for starting therapy. In particular, the absolute lymphocyte count was not used as the sole indicator for treatment. Active disease, requiring therapy, was defined when at least one of the criteria set out in the NCI-sponsored Working Group guidelines was satisfied).

Additional outcome(s)

· not applicable

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 337 participants, 91 events

Which model was used?

· Authors used nomogram/formula

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

no

PROBAST: Applicability

Not applicable, outcome did not match primary outcome of the model.

Notes

Funding & conflict of interest

- "This work was supported by funding from Associazione Italiana per la Ricerca sul Cancro (AIRC 5xmille grant 9980, IG10492 to MF and FM and IG10136 to AN). We thank AIL Cosenza-Fondazione Amelia Scorza' onlus, Cosenza, Italy, and Brigida Gulino for precious secretarial assistance." (Gentile 2016)
- Conflict of interest not reported

Other comments

Cohort identical in several publications



MDACC 2007 V - Muñoz-Novas 2018 (Spanish cohort)

Study characteristics

General information

Model and type of study

- Munoz-Novas 2018 (Spanish cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

• median 46 months, range not reported

Participants

Number of included participants in the cohort

· 696 participants

Setting

· Spain, multicentre study

Recruitment period

• 1989 - 2013

Age (in years)

• median 67 years (range: 25 - 90 years)

Sex

• 64.2 % male

Stages of disease

- Rai 0-II: 93.8%; Rai III-IV: 6.2%
- Binet A/B/C: not reported

Treatment

- 186 received treatment
- · unclear which treatment

Inclusion criteria

• all participants diagnosed with CLL

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



MDACC 2007 V - Muñoz-Novas 2018 (Spanish cohort) (Continued)

Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)

Timing of predictor measurement

· at diagnosis, data from medical records

Outcome(s)

Primary outcome in study

 overall survival: overall survival (OS) was calculated from the time of diagnosis to death or last follow-up.

Additional outcome(s)

• time-to-first-treatment: time-to-first-therapy (TTFT) from the date of diagnosis to first treatment or last follow-up

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

· yes (213 participants)

If yes, how was missing data handled?

complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

- overall survival: 483 participants, 92 events
- time-to-first-treatment: 483 participants, 186 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (1-3);(4-7);(above 7))
- at 5 years
- percentage of group without event at time point (low risk: 98.4%, intermediate risk: 79.6%, high risk: 51.1%)

PROBAST: Applicability

<u>Domain 1: Participant selection</u>

low

Domain 2: Predictors

low

Domain 3: Outcome



MDACC 2007 V - Muñoz-Novas 2018 (Spanish cohort) (Continued)

Iov

Notes

Funding & conflict of interest

- · Funding not reported
- "The authors declare that there are no conflicts of interest regarding the publication of this article."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Retrospective cohort: "A total of 696 unselected CLL patients newly diagnosed and previously untreated from different institutions of the central region of Spain were included in this study." Patients with missing values were later excluded at analysis stage, rated in domain 4.
Domain 2: Predictors	Yes	Routine clinical data assessed at diagnosis was used (from medical records).
Domain 3: Outcome	Yes	Objective standard outcome: no definition of outcome and assessment - we assumed that definition and assessment of the objective outcome OS was similar and therefore did not rate as high risk.
Domain 4: Analysis	No	Low number of event; exclusion of patients with missing values from analysis; calibration not reported
Overall judgement	No	

MDACC 2007 V - Pflug 2014 (3 RCTs)

Study characteristics

General information

Model and type of study

- Pflug 2014 (German CLL study group cohort)
- Validation study

Secondary citations

· not applicable

Language of publication

English

Study design

• combined data (e.g. IPD or combination of cohorts)

Follow-up time

• median 63.4 months, range not reported

Participants

Number of included participants in the cohort

• 2007 participants



MDACC 2007 V - Pflug 2014 (3 RCTs) (Continued)

Setting

 several, multicentre study (Australia, Austria, Belgium, Czech Republic, Denmark, France, Germany, Israel, Italy, New Zealand, Spain)

Recruitment period

1997 - 2006

Age (in years)

• median 60 years (range: 30 - 81 years)

Sex

• 68.4% male

Stages of disease

- Rai 0: 20.8%; Rai I: 18.4%; Rai II: 37.9%; Rai III: 8.6%; Rai IV: 14.3%
- Binet A: 42.6%; Binet B: 37.7%; Binet C: 19.8%

Treatment

- · participants were treated with:
- early fludarabine (F)
- F plus cyclophosphamide (FC)
- FC plus rituximab (FCR)

Inclusion criteria

- CLL1 trial: 18-75 years, confirmed diagnosis of B-cell chronic lymphocytic leukaemia, Rai stage 0-II
 (Binet stage A), life expectancy above 6 months, no autoimmune haemolytic anaemia, no thrombocytopenia, no severe organ dysfunction, no other prior or concurrent malignancy, no prior or other concurrent chemotherapy
- CLL4 trial: < 65 years with untreated CLL in advanced stage (all Binet stage C patients; Binet stage B with symptoms, which require therapy; Binet stage A with severe B-symptoms)
- CLL8 trial: >= 18 years, diagnosed B-cell chronic lymphocytic leukaemia (CLL) defined by the National
 Cancer Institute (NCI) Working Group criteria, Binet stage C disease OR Binet stage B disease AND 1 of
 other predefined signs or symptoms, ECOG performance status 0-1, Cumulative Illness Rating Scale
 (CIRS) score above 6, life expectancy above 6 months, no active infection, no pregnancy or nursing,
 no concurrent or previous treatment of CLL by chemotherapy, radiotherapy, or immunotherapy, and
 more

Exclusion criteria

 CLL4 trial: age < 18 and > 65 years were excluded as well as patients with any previous treatment of CLL, life expectancy less than six months and an Eastern Cooperative Oncology Group performance status of more than two. Patients were also excluded if they had severe organ dysfunction, concomitant or previous other neoplasms or an autoimmune haemolytic anaemia or thrombocytopenia.

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)
- no changes

Timing of predictor measurement

• at study entry (< 1 year: 63.3%, > 1 year: 36.7%)



MDACC 2007 V - Pflug 2014 (3 RCTs) (Continued)

Outcome(s)

Primary outcome in study

overall survival: OS was defined as the time between registration/randomisation and death.

Additional outcome(s)

• progression-free survival: PFS was calculated from registration/randomisation to start of the first CLL treatment or disease progression or death.

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

• yes (863 participants)

If yes, how was missing data handled?

· complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

overall survival: 1144 participants, number of events not reported in publication - 180 at 5 years

Which model was used?

· c-statistic reported for the nomogram

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

no

PROBAST: Applicability

Domain 1: Participant selection

• unclear - restricted inclusion and exclusion criteria based on RCTs

Domain 2: Predictors

low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- "This manuscript was written on behalf of the German CLL Study Group. Studies CLL1, CLL4, and CLL8 were planned and conducted as investigator-initiated trials by the German CLL Study Group and were supported by research grants from German Cancer Aid, Medac Schering Onkologie, and F. Hoffmann-La Roche. T.D.S. is a clinical scholar of the Leukemia Lymphoma Society."
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MDACC 2007 V - Pflug 2014 (3 RCTs) (Continued)

Mundipharma, and received honoraria and research funding from both. H.D. received research grants from Roche. U.J. received honoraria and research funding from Roche. M.H. is a consultant and/or holds an advisory role and received research funding from Roche. J.B., M.A.B., T.E., K.B., G.M., K.G.R., M.J.E., G.H., R.B., A.-M.F., C.-M.W., K.F., and N.E.K. declare no competing financial interests."

Other comments

• The authors provided additional data upon request (classification tables).

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Overlap of individuals between trials prevented; only concern could be comparability of the three trials.
Domain 2: Predictors	Yes	Prospective collection of predictors within clinical trials
Domain 3: Outcome	Yes	Objective standard outcome: follow-up in frame of each RCT
Domain 4: Analysis	No	Number of events sufficient; individuals with missing data were excluded from model development. It was not clear if this was the case for the validation of the MDACC2007 model; calibration not reported
Overall judgement	No	

MDACC 2007 V - Rani 2018	3 (Indian cohort)		
Study characteristics			
General information	Model and type of study		
	Rani 2018 (Indian cohort)Validation study		
	<u>Secondary citations</u>		
	not applicable		
	Language of publication		
	• English		
	Study design		
	• unclear		
	Follow-up time		
	median 40.5 months (range not reported)		
Participants	Number of included participants in the cohort		
	• 198 participants		
	Setting		
	India, unclear if single or multicentre study		
	Recruitment period		



MDACC 2007 V - Rani 2018 (Indian cohort) (Continued)

not reported

Age (in years)

median 60 years (range: not reported)

Sex

• 77% male

Stages of disease

- Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%
- · Binet A/B/C: not reported

Treatment

- 62 received chlorambucil-based therapy
- 56 received rituximab-based therapy
- 20 received other therapies

Inclusion criteria

• treatment-naive CLL patients according to Hallek 2008

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)

Timing of predictor measurement

· not reported

Outcome(s)

Primary outcome in study

• overall survival: OS was defined as the time from the date of diagnosis to date of death or date of last follow-up.

Additional outcome(s)

• not applicable for this model (TTFT was assessed in this cohort for other prognostic models)

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• overall survival: 198 participants, 86 events



MDACC 2007 V - Rani 2018 (Indian cohort) (Continued)

Which model was used?

• simplified model

Was the model updated?

• nc

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups
- · survival probabilities not reported

PROBAST: Applicability

Domain 1: Participant selection

· unclear - recruitment period and eligibility criteria unclear

Domain 2: Predictors

• unclear - no information on timing of predictor because the study design was unclear

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "The financial support was provided by the Department of Biotechnology (BT/PR11106/ GBD/27/145/2008, BT/PR15438/MED/30/606/2011 and T/PR8680/AGR/36/754/2013), Ministry of Science and Technology, GOI, and All India Institute of Medical Sciences, New Delhi (8-60/A060/ 2011/ RS) to RG for carrying out this work."
- "The authors declare that they have no conflict of interest."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	No clear eligibility criteria; possibly missing values as exclusion reason; no recruitment period
Domain 2: Predictors	Yes	Study design unclear, " were obtained from medical records of the patients". Routine clinical predictors were used in this model, therefore we rated this study as low.
Domain 3: Outcome	Unclear	Short follow-up time
Domain 4: Analysis	No	Low number of events; calibration not reported; survival per risk groups graphically presented
Overall judgement	No	



MDACC 2007 V - Trajkova 2013 (Macedonia)

Study characteristics

General information

Model and type of study

- Trajkova 2013 (Macedonian cohort)
- · Validation study

Secondary citations

· not applicable

Language of publication

· English

Study design

unclear

Follow-up time

· not reported

Participants

Number of included participants in the cohort

· 100 participants

Setting

• Macedonia, single-centre study

Recruitment period

• 2011 - 2013

Age (in years)

• median 64.8 years (range: 47 - 78 years)

Sex

• 63% male

Stages of disease

- Rai 0: 54%; Rai I: 5%; Rai II: 32%; Rai III: 5%; Rai IV: 4%
- Binet A/B/C: not reported

Treatment

- 41 received treatment
- · unclear which therapy

Inclusion criteria

• Our study incorporated 100 consecutive treatment-naive CLL patients with IgHV mutational status.

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>



MDACC 2007 V - Trajkova 2013 (Macedonia) (Continued)

Age (continuous: no transformation; in years); ALC (absolute lymphocyte count) (continuous: no transformation; x10^9/L); serum ß2-microglobulin (continuous: no transformation); nodal groups (binary); Rai stage (categorical; binary: stage III or IV vs. 0-II); gender (binary)

Timing of predictor measurement

• at presentation at University Haematology Clinic

Outcome(s)

Primary outcome in study

time-to-first-treatment: time-to-event end point was defined as the time from first visit to the University Haematology Clinic to first CLL treatment. There was no restriction of time from diagnosis to presentation at the University Haematology Clinic.

Additional outcome(s)

not applicable

Outcome in model development

· overall survival

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 100 participants, 41 events

Which model was used?

unclear

Was the model updated?

unclear

Performance measures reported?

- Calibration: not reported
- Discrimination: not reported

Creation of risk groups?

- ves
- No reporting of observed survival per subgroup

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest



MDACC 2007 V - Trajkova 2013 (Macedonia) (Continued)

not reported

Other comments

• "estimated and projected" survival without any comparison of observed survival provided; we could not use this information for our review and therefore, we did not continue with data extraction.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	"100 consecutive treatment naïve CLL patients"
Domain 2: Predictors	Yes	Routine clinical data
Domain 3: Outcome	Unclear	No observation time reported; outcome assessment not described
Domain 4: Analysis	No	Low number of participants and events; not really validating MDACC 2007 - reporting of projected survival only and additional univariable analysis
Overall judgement	No	

MDACC 2011 D - Wierda 2011 (MDACC)

Study characteristics		
General information	Model and type of study	
	 Wierda 2011 (development (MDACC 2004-2009)) Development study 	
	<u>Secondary citations</u>	
	not applicable	
	Language of publication	
	• English	
	Study design	
	retrospective cohort	
	Follow-up time	
	• median 26 months (range 3 - 73 months)	
Participants	Number of included participants in the cohort	

• US (Texas), single-centre study

Recruitment period

• 930 participants

• 2004 - 2009



MDACC 2011 D - Wierda 2011 (MDACC) (Continued)

Age (in years)

• median 59 years (range: 30 - 89 years)

Sex

• 61% male

Stages of disease

• Rai 0: 36%; Rai I: 51%; Rai II: 8%; Rai III-IV: 5%

Treatment

- 232 received treatment
- · unclear which treatment

Inclusion criteria

• no previous treatment; not recommended for first-line treatment at initial visit

Exclusion criteria

· not reported

Predictors

Number of candidate predictors

- univariable analysis: more than 25
- · multivariable analysis: unclear

List of predictors in final model (including cut-points for dichotomised factors)

IgHV status (continuous: categorical/binary transformation); diameter of largest palpated cervical LN, cm (continuous: no transformation); FISH category (categorical; del(11q) or del(17p) vs. none); number of involved lymph node sites (continuous: categorical/binary transformation); lactate dehydrogenase IU/L/100 (continuous: no transformation); IgHV status*LDH (interaction term)

Timing of predictor measurement

at study entry (median time from diagnosis to presentation to MDACC was 3.4 months, range 0 to 428 months)

Outcome(s)

Primary outcome in study

 time-to-first-treatment: TTFT was defined as the time interval between the date of presentation to MDACC and date of first CLL treatment.

Additional outcome(s)

not applicable

Missing data

Participants with any missing data?

• yes (243 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 687 participants, 193 events

Predictor selection method



MDACC 2011 D - Wierda 2011 (MDACC) (Continued)

• univariable analysis, during multivariable modelling: stepwise regression mix, (P value or CI)

Statistical method

• Cox proportional hazard model

Simplification of model?

yes

Performance measures reported?

- Calibration: not reported
- Discrimination: not reported

Creation of risk groups?

no

PROBAST: Applicability

Domain 1: Participant selection

• unclear - individuals with missing values were excluded. We were unsure if the sample was representative for all individuals with CLL

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "Supported in part by the Leukemia and Lymphoma Society Clinical Scholar program (W.G.W.)"
- "The author(s) indicated no potential conflicts of interest."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant selection	No	Individuals without evaluation of clinical factors and at least one newer factor were excluded: "We identified 930 previously untreated patients who presented to MD Anderson Cancer Center (MDACC) between January 2004 and December 2009, were not recommended for first-line treatment at initial visit, and were evaluated for traditional clinical and laboratory prognostic factors and one or more of the newer prognostic factors including IgHV mutation status, chromosome abnormalities by FISH analysis, and ZAP-70 expression by flow cytometry and/or immunohistochemistry (IHC)."
Domain 2: Predictors	Yes	Predictor assessment well described; comparison of assessment tools
Domain 3: Outcome	No	Some univariably tested predictors were not excluded from the outcome definition (progressive or symptomatic splenomegaly, massive nodes, progressive or symptomatic lymphadenopathy, Rai stage; Hallek 2008); short observation time
Domain 4: Analysis	No	Many predictors compared to events, 193 events (more than 25 predictors); univariable selection of predictors; unclear how many predictors entered in-



MDACC 2011 D - Wierda 2011 (MDACC) (Continued)

to MV analysis; no performance measures reported; split sample, but after the model was already developed, which means that the data were internally validated on data that it was developed with. Bootstrapping was done, however not used to correct the model; assumption of proportionality was checked

Overall judgement

No

MDACC 2011 V - Molica 2016 (O-CLL1-GISL)

Study characteristics

General information

Model and type of study

- Molica 2013 (O-CLL1-GISL)
- · Validation study

Secondary citations

Molica 2013; Molica 2014; Molica 2016; Molica 2017; Molica 2017

Language of publication

English

Study design

· prospective cohort

Follow-up time

• median 30 months (range 1 - 65 months)

Participants

Number of included participants in the cohort

• 328 participants

<u>Setting</u>

· Italy, multicentre study

Recruitment period

• 2006 - 2010

Age (in years)

• median 61 years (range: 33 - 70 years)

<u>Sex</u>

• 58.2% male

Stages of disease

- Rai 0: 76.5%; Rai I-II: 23.5%
- Binet A: 100%

Treatment

- 91 received treatment
- · unclear which treatment



MDACC 2011 V - Molica 2016 (O-CLL1-GISL) (Continued)

Inclusion criteria

The inclusion criteria for CLL diagnosis, which were used at the time of study design and initiation, followed the National Cancer Institute-sponsored Working Group guidelines, which require absolute lymphocytosis with a lower threshold of above 5000 mature-appearing lymphocytes/mL in the peripheral blood. Because the objective of this observational study was to evaluate the role of novel prognostic variables in younger patients with CLL, only those aged above 70 years were considered eligible.

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

IgHV status (continuous: categorical/binary transformation); diameter of largest palpated cervical LN, cm (continuous: no transformation); FISH category (categorical; del(11q) or del(17p) vs. None); number of involved lymph node sites (continuous: categorical/binary transformation); lactate dehydrogenase IU/L/100 (continuous: no transformation); IgHV status*LDH (interaction term)

Timing of predictor measurement

· at study entry, within 12 months of diagnosis

Outcome(s)

Primary outcome in study

time-to-first-treatment: TTFT was defined as the interval between the date of registration and the date
of initiation of first CLL treatment. Patients underwent sequential monitoring, and the frequency of
follow-up visits was individualised according to patient risk; this ranged from 3 to 6 months (median
6 months). All physicians who registered patients in this observational database stated that they had
used the NCI-WG guidelines as a reference criterion for starting therapy. In particular, the absolute
lymphocyte count was not used as the sole indicator for treatment. Active disease, requiring therapy,
was defined when at least one of the criteria set out in the NCI-sponsored Working Group guidelines
was satisfied.

Additional outcome(s)

· not applicable

Outcome in model development

· time-to-first-treatment

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- time-to-first-treatment: 337 participants, 91 events
- in Molica 2013: time-to-first-treatment: 328 participants, 68 events

Which model was used?

Cox proportional hazard model

Was the model updated?

no



MDACC 2011 V - Molica 2016 (O-CLL1-GISL) (Continued)

Performance measures reported?

Calibration: reported (calibration plot)

• Discrimination: reported (c-statistic)

Creation of risk groups?

no

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "This work was supported by grants from Associazione Italiana Ricerca sul Cancro (AIRC) (to Antonino Neri -IG4569, MFIG10492 and Fortunato Morabito -RG6432) and AIRC-Special Program Molecular Clinical Oncology-"5 per mille", grant 9980, 2010-15 to Antonino Neri, Manlio Ferrarini and Fortunato Morabito; Ricerca Finalizzata from Italian Ministry of Health 2006 (GC, FM and MF) and 2007 (GC)."
- "The authors made no disclosures."

Other comments

• The O-CLL1-GISL cohort was used in various publications and to validate different CLL models.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Prospective cohort, detailed eligibility criteria in trial registry
Domain 2: Predictors	Yes	Predictors prospectively collected
Domain 3: Outcome	No	See development of the model: some predictors in the model were not excluded from the outcome definition (number and size of lymph nodes included in model, however these factors also formed Binet stage, which was a reason for treatment indication, Hallek 2008).
Domain 4: Analysis	No	Low number of events; no information on missing data reported
Overall judgement	No	

Morabito D - Morabito 2009 (Italian cohort)

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Studv	cna	racte	ristics

General information

Model and type of study

- Morabito 2009 (Development cohort (Italian))
- Development study



Morabito D - Morabito 2009 (Italian cohort) (Continued)

Secondary citations

· not applicable

Language of publication

• English

Study design

· not reported

Follow-up time

• median 36 months (range 12 - 180 months)

Participants

Number of included participants in the cohort

· 262 participants

Setting

· Italy, multicentre study

Recruitment period

· not reported

Age (in years)

• median 65 years (range not reported)

Sex

• 61.5% male

Stages of disease

- Rai 0-IV: not reported
- Binet A: 100%

Treatment

· not reported

Inclusion criteria

 diagnosis of typical CLL based on the National Cancer Institute (NCI) Working Group criteria (Hallek 2008) and confirmed by flow cytometry

Exclusion criteria

not reported

Predictors

Number of candidate predictors

- univariable analysis: 3
- multivariable analysis: not applicable

<u>List of predictors in final model (including cut-points for dichotomised factors)</u>

• IgHV mutational status (continuous: categorical/binary transformation; 2% threshold, as established in literature); CD38 (continuous: categorical/binary transformation; 30% threshold); ZAP-70 (continuous: categorical/binary transformation; threshold established by ROC curve)

Timing of predictor measurement



Morabito D - Morabito 2009 (Italian cohort) (Continued)

 markers were tested within two years in 70% of cases, within 4 years in 15% and from 4 to 5 years from diagnosis in the remaining 15%

Outcome(s)

Primary outcome in study

time-to-first-treatment: TTT was measured from diagnosis to first-line treatment or last follow-up.
 Treatment was decided uniformly in all participating centres based on documented progressive and symptomatic disease according to NCI working guidelines.

Additional outcome(s)

not applicable

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 252 participants, 51 events

Predictor selection method

univariable analysis only (P value or CI); a scoring system was designed in which one point was assigned to each unfavourable prognostic marker and the final score was determined by the sum of unfavourable markers present.

Statistical method

• univariable analysis: Kaplan-Meier method

Simplification of model?

not applicable

Performance measures reported?

- Calibration: not reported
- · Discrimination: not reported

Creation of risk groups?

no

PROBAST: Applicability

Domain 1: Participant selection

unclear - not sufficient information (eligibility criteria, sample characteristics, inclusions and exclusions) on the sample to make a judgement

Domain 2: Predictors

• high - predictors not collected at study entry or at diagnosis: "Markers were tested within two years in 70% of cases, within 4 years in 15% and from 4 to 5 years from diagnosis in the remaining 15%."

Domain 3: Outcome

• low

Notes

Funding & conflict of interest



Morabito D - Morabito 2009 (Italian cohort) (Continued)

- "Supported in part from Associazione Italiana Ricerca sul Cancro (AIRC) (to FM, AN and MF), FIRB (Grant no RBIP06LCA9, to MF) MIUR, CIPE (2006–8, CBA project, to SZ), ISS (to SZ), Progetti Strategici Ricerca Finalizzata Ministero Italiano della Salute 'RFPS_2006_3_33_99_60' (to GC) and 'RF-PS_2006_340196' (to FM and MF); progetto ordinario ricerca finalizzata Ministero Italiano della salute-2007 (to GC), progetto Compagnia San Paolo (to GC), the Fondazione Internazionale Ricerche Medicina Sperimentale (FIRMS) provided financial and administrative assistance, Fondazione 'Amelia Scorza' onlus, Cosenza, Italy; and Associazione Italiana contro le Leucemie -Milano. KT and SM are supported by fellowships from the Fondazione Italiana Ricerca sul Cancro (FIRC)".
- · Conflicts of interests not reported

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Only one statement on inclusion criteria. No statement on baseline sample and possible exclusions. Very little baseline information on participants
Domain 2: Predictors	Yes	Prospective collection of predictor information, centralised laboratory
Domain 3: Outcome	Yes	Standard outcome definition used: "Treatment was decided uniformly in all participating centres based on documented progressive and symptomatic disease according to NCI working guidelines".
Domain 4: Analysis	No	Not sufficient events; continuous data not handled appropriately (ROC to find the 'best' cut-point); no information on missing data; no performance measure reported
Overall judgement	No	

Morabito V - Gentile 2014 (O-CLL1-GISL)

Study characteristics

General information	Model and to

Model and type of study

- Gentile 2014 (O-CLL1-GISL)
- Validation study

Secondary citations

· not applicable

Language of publication

English

Study design

· prospective cohort

Follow-up time

median 38.5 months (range 6 - 82 months)

Participants <u>Number of included participants in the cohort</u>



Morabito V - Gentile 2014 (O-CLL1-GISL) (Continued)

480 participants

Setting

· Italy, multicentre study

Recruitment period

· 2007 - not reported

Age (in years)

• 47.4 % of individuals were aged < 60 years

Sex

• 58.5% male

Stages of disease

- Rai 0: 77.3%; Rai I-II: 22.7%
- Binet A: 100%

Treatment

· not reported

Inclusion criteria

 the inclusion criteria for CLL diagnosis, employed at the time of study start, followed the NCI/WG guidelines established in 1996, required > 5.000 lymphocytes/mL in the peripheral blood.

Exclusion criteria

exclusion criteria were the following: (i) CD5- and/or CD23- B-lymphoproliferative disorders; (ii) clinical Binet stage B or C, and (iii) need of therapy according to NCI/WG guidelines; (iv) age > 70 years.

Predictors

List of predictors in final model (including cut-points for dichotomised factors)

- IgHV mutational status (continuous: categorical/binary transformation; 2% treshold, as established in literature); CD38 (continuous: categorical/binary transformation; 30% treshold); ZAP-70 (continuous: categorical/binary transformation; threshold established by ROC curve)
- no difference between development and validation cohort predictors

Timing of predictor measurement

• at study entry, maximum 1 year after diagnosis

Outcome(s)

Primary outcome in study

• progression-free survival: PFS has been defined as time to therapy requirement.

Additional outcome(s)

· not applicable

Outcome in model development

• time-to-first-treatment

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?



Morabito V - Gentile 2014 (O-CLL1-GISL) (Continued)

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• progression-free survival: 468 participants, 121 events

Which model was used?

· original model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- · Discrimination: not reported

Creation of risk groups?

- yes, 3 risk groups, (cut-points at 1; 2-3; 4)
- · at 3 years
- percentage of group without event at time point (low risk: 91.7%, intermediate risk: 82.9%, high risk: 57.4%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

• low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- "Associazione Italiana per la Ricerca sul Cancro; Contract grant no.: 9980, IG10492, IG10136"
- "Nothing to report"

Other comments

- The O-CLL1-GISL cohort was used in various publications and to validate different CLL models.
- The corresponding author provided additional information regarding the cohort.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Prospectively registered study (O-CLL1-GISL) with defined inclusion and exclusion criteria
Domain 2: Predictors	Yes	Prospective collection of predictor information. Predictors assessed study entry at a maximum of 1 year after diagnosis.
Domain 3: Outcome	Yes	CLL progression criteria 2008 were used. Outcome labelled differently than model development study. However, progression-free survival (PFS) has been defined as time-to-therapy requirement, which is equivalent to time-to-treatment in Morabito 2009.



Morabito V - Gentile 2014 (O-CLL1-GISL) (Continued)

Domain 4: Analysis

Unclear

No relevant performance measures reported (progression per risk group was

reported)

O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL)

Study characteristics

General information

Model and type of study

- Gentile 2016 (O-CLL1-GISL)
- Development study

Secondary citations

• Molica 2016; Molica 2017; Molica 2017

Language of publication

· English

Study design

· prospective cohort

Follow-up time

• median 42 months (range 6 - 82 months)

Participants

Number of included participants in the cohort

· 480 participants

Setting

• Italy, multicentre study

Recruitment period

• 2007 - not reported

Age (in years)

• 46.7% of individuals were younger than 60 years

<u>Sex</u>

• 57.3% male

Stages of disease

- Rai 0: 77.7%; Rai I-II: 22.3%
- Binet A: 100%

Treatment

- 84 out of 337 received treatment
- · unclear which treatment

Inclusion criteria

• Established diagnosis of B-CLL by NCI criteria, performed by local haematologist (confirmed by the biological review committee according to flow cytometry analysis (positive clusters of differentiation



O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL) (Continued)

antigen 5 (CD5), 19 (CD19), 23 (CD23)). Age more than 18 years and less than 70 years. Eastern Cooperative Oncology Group (ECOG) less than or equal to 2. Binet stage A. Diagnosis performed within 12 months before inclusion in the study. NCI watch-and-wait policy.

Exclusion criteria

Patients with CLL whose diagnosis exceeded 12 months before registration. Patients with leukaemic
phase of lymphoproliferative disorders of B origin CD5- and/or CD23- according to flow cytometry
analysis. Clinical Binet stage B or C. Patients who necessitated therapy according to NCI guidelines.
Age > 70 years. Without a written informed consent.

Predictors

Number of candidate predictors

- univariable analysis: 10
- multivariable analysis: 7

List of predictors in final model (including cut-points for dichotomised factors)

Rai stage (categorical; 0 vs. I/II); ALC (continuous: categorical/binary transformation); beta2-mi-croglobulin (continuous: categorical/binary transformation; normal/elevated (elevated above 2 mg/L)); IgHV status (continuous: categorical/binary transformation)

Timing of predictor measurement

at study entry (< 12 months from diagnosis)

Outcome(s)

Primary outcome in study

 time-to-first-treatment: TFS was calculated from diagnosis to the first CLL treatment (event) or to last follow-up (censoring)

Additional outcome(s)

not applicable

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 337 participants, 84 events

Predictor selection method

• univariable analysis, unclear if stepwise approach was used in multivariable modelling (criterion: P value or CI)

Statistical method

· Cox proportional hazard model

Simplification of model?

yes

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)



O-CLL-1 D - Gentile 2016 (O-CLL-1-GISL) (Continued)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (0-2); (3-5); (6-7))
- at 3 years
- percentage of group without event at time point (low risk: 95.3%, intermediate risk: 74.5%, high risk: 28.6%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "Associazione Italiana per la Ricerca sul Cancro (AIRC 5xmille grant 9980, IG10492 to MF and FM and IG10136 to AN)."
- "The authors declare no conflict of interest."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Prospective cohort, clear inclusion and exclusion criteria in trial registry
Domain 2: Predictors	Yes	Prospective and standardised collection of predictor information ("CLL cell phenotype, CD38 and ZAP-70 expression, and <i>IgHV</i> mutational status were performed in a central laboratory in Genova, while all FISH and genetic analyses were performed in Milan").
Domain 3: Outcome	No	Standard outcome definition, however, predictors were not excluded from outcome definition (e.g. Rai stage).
Domain 4: Analysis	No	Continuous data not handled appropriately ("Continuous variables of prognostic importance on TFS in univariate proportional hazards Cox regression were dichotomized using published thresholds and laboratory norms").
Overall judgement	No	

O-CLL-1 V - Rani 2018 (Indian cohort)

Study characteristics

General information

Model and type of study

- Rani 2018 (Indian cohort)
- · Validation study



O-CLL-1 V - Rani 2018 (Indian cohort) (Continued)

Secondary citations

· not applicable

Language of publication

• English

Study design

unclear

Follow-up time

• median 40.5 months (range: 1 - 215 months)

Participants

Number of included participants in the cohort

· 198 participants

Setting

• India, unclear if single or multicentre study

Recruitment period

· not reported

Age (in years)

• median 60 years (range not reported)

Sex

• 77% male

Stages of disease

- Rai 0: 14.6%; Rai I: 21.2%; Rai II: 34.9%; Rai III: 14.1%; Rai IV: 15.2%
- Binet A/B/C: not reported

Treatment

- 62 received chlorambucil-based therapy
- · 56 received rituximab-based therapy
- 20 received other therapies

Inclusion criteria

• treatment-naive CLL participants according to Hallek 2008

Exclusion criteria

not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

gender (binary); age (continuous, dichotomisation after univariable analysis); ECOG PS (categorical, dichotomisation after univariable analysis); del(17p) (continuous: categorical/binary transformation); del(11q) (continuous: categorical/binary transformation; literature-based); IgHV mutational status (continuous: categorical/binary transformation; literature-based); s-TK (continuous, i after univariable analysis); s-beta2m (continuous, dichotomisation after univariable analysis)

Timing of predictor measurement



O-CLL-1 V - Rani 2018 (Indian cohort) (Continued)

· not reported

Outcome(s)

Primary outcome in study

• time-to-first-treatment: TTFT was defined as the time from the date of diagnosis to date of commencement of first therapy.

Additional outcome(s)

· not applicable

Outcome in model development

· time-to-first-treatment

Missing data

Participants with any missing data?

no

If yes, how was missing data handled?

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 140 participants, 89 events

Which model was used?

· simplified model

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- · Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (0-2); (3-5); (6-7))
- median survival (low risk: 3 years, intermediate risk: 24 years, high risk: 79 years)

PROBAST: Applicability

Domain 1: Participant selection

• unclear - recruitment period unclear

Domain 2: Predictors

• unclear - no information on timing of predictor because the study design was unclear

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "The financial support was provided by the Department of Biotechnology (BT/PR11106/ GBD/27/145/2008, BT/PR15438/MED/30/606/2011 and T/PR8680/AGR/36/754/2013), Ministry of Science and Technology, GOI, and All India Institute of Medical Sciences, New Delhi (8-60/A060/2011/ RS) to RG for carrying out this work."
- "The authors declare that they have no conflict of interest."



O-CLL-1 V - Rani 2018 (Indian cohort) (Continued)

Other comments

• Graphs were provided to obtain the percentage of participants untreated at a specific time point, however, these graphs were printed very small.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Inclusion and exclusion criteria unclear; it seemed to us that only individuals with complete information on all predictors were included; recruitment period unclear; study design unclear
Domain 2: Predictors	Yes	Not explicitly stated, but predictors probably assessed in a similar way (single-centre)
Domain 3: Outcome	No	See model development: Predictors were not excluded from outcome definition (e.g. Rai stage).
Domain 4: Analysis	No	Low number of events; no information on handling of missing values
Overall judgement	No	

O-CLL1 V - Gentile 2016 (Mayo cohort)

Study o	charact	eristics
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General	info	rmati∧n
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Model and type of study

- Gentile 2016 (Mayo cohort 2001 2008)
- · Validation study

Secondary citations

· not applicable

Language of publication

• English

Study design

not reported

Follow-up time

• median 97 months (range not reported)

Participants

Number of included participants in the cohort

• 428 participants

Setting

• USA, single-centre study

Recruitment period

• 2001 - 2008



O-CLL1 V - Gentile 2016 (Mayo cohort) (Continued)

Age (in years)

• 40.2% <= 60 years

Sex

• 66.4% male

Stages of disease

- Rai 0: 67.5%; Rai I-II: 32.5%
- Binet A/B/C: not reported

Treatment

- · 130 were treated
- · unclear which treatment

Inclusion criteria

· not reported

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

Rai stage (categorical; 0 vs. I/II); ALC (continuous: categorical/binary transformation); beta2-microglobulin (continuous: categorical/binary transformation; normal/elevated (elevated above 2 mg/L)); IgHV status (continuous: categorical/binary transformation)

Timing of predictor measurement

• within 18 months of diagnosis

Outcome(s)

Primary outcome in study

 time-to-first-treatment: TFS was calculated from diagnosis to the first CLL treatment (event) or to last follow-up (censoring).

Additional outcome(s)

• not applicable

Outcome in model development

· time-to-first-treatment

Missing data

Participants with any missing data?

· not reported

If yes, how was missing data handled?

· not reported

Analysis

Number of participants & number of events (specific time points where reported)

• time-to-first-treatment: 428 participants, 130 events

Which model was used?

· simplified model



O-CLL1 V - Gentile 2016 (Mayo cohort) (Continued)

Was the model updated?

no

Performance measures reported?

- Calibration: yes (Hosmer-Lemeshow test)
- · Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 3 risk groups, (cut-points at (0-2); (3-5); (6-7))
- · at 3 years
- percentage of group without event at time point (low risk: 95.5%, intermediate risk: 78.9%, high risk: 40.6%)

PROBAST: Applicability

Domain 1: Participant selection

· unclear - recruitment period unclear

Domain 2: Predictors

• low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- · "Associazione Italiana per la Ricerca sul Cancro (AIRC 5xmille grant 9980, IG10492 to MF and FM and IG10136 to AN)."
- "The authors declare no conflict of interest."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	No eligibility criteria; individuals with missing values were excluded.
Domain 2: Predictors	Yes	Not explicitly stated, but predictors probably assessed in a similar way (single-centre).
Domain 3: Outcome	No	See model development: Predictors not excluded from outcome definition (e.g. Rai stage)
Domain 4: Analysis	Unclear	Calibration not reported; individuals with missing values were probably excluded from analysis, rated in domain 1.
Overall judgement	No	



Rossi D - Rossi 2013 (Italian cohort)

Study characteristics

General information

Model and type of study

- Rossi 2013
- Development study

Secondary citations

· not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

• median 67.2 months (range not reported)

Participants

Number of included participants in the cohort

• 637 participants

Setting

· Italy, multicentre study

Recruitment period

• 1996 - 2011

Age (in years)

• 51.2% of individuals were older than 65 years.

Sex

• 58.4% male

Stages of disease

- Rai 0: 74.9%; Rai I-II: 11.5%
- · Binet A: not reported

Treatment

- 122 patients received rituximab-based regimens (i.e. fludarabine-cyclophosphamide-rituximab, fludarabinerituximab, or pentostatin-cyclophosphamide-rituximab)
- 64 received fludarabine-based regimens (i.e. fludarabine or fludarabinecyclophosphamide)
- 80 received alkylator-based regimens (i.e. chlorambucil)

Inclusion criteria

• 'newly diagnosed and previously untreated CLL patients of whom 583 (91.5%) were provided with regular follow-up'

Exclusion criteria

not reported

Predictors

Number of candidate predictors



Rossi D - Rossi 2013 (Italian cohort) (Continued)

- · univariable analysis: no univariable analysis
- multivariable analysis: 18

<u>List of predictors in final model (including cut-points for dichotomised factors)</u>

• TP53 (binary); BIRC3 DIS (binary); SF3B1 M (binary); NOTCH1 M (binary); del11q22-q23 (binary)

Timing of predictor measurement

· at diagnosis

Outcome(s)

Primary outcome in study

• overall survival: OS was measured from the date of initial presentation to the date of death from any cause (event) or last follow-up (censoring).

Additional outcome(s)

 Treatment-free survival: TFS was measured from date of initial presentation to date of progressive disease requiring treatment according to IWCLL-NCI guidelines (event), death or last follow-up (censoring).

Missing data

Participants with any missing data?

• yes (54 participants)

If yes, how was missing data handled?

· not reported

Analysis

Number of participants & number of events (specific time points where reported)

• overall survival: 583 participants, 178 events

Predictor selection method

• full model, during multivariable modelling: hierarchical clustering

Statistical method

recursive partitioning

Simplification of model?

no

Performance measures reported?

- · Calibration: not reported
- · Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups, (cut-points at very low risk: del13q14 sole lesion; low risk: +12 or all lesions; intermediate: NOTCH1 and/or SF3B1; high risk: TP53 and/or BIRC3)
- · at 5 years
- percentage of group without event at time point (low risk: 86.9%, intermediate risk: 77.6%, high risk: 65.9%, very high risk: 50.9%)

PROBAST: Applicability

Domain 1: Participant selection

• low

Domain 2: Predictors



Rossi D - Rossi 2013 (Italian cohort) (Continued)

low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- "This study was supported by the Associazione Italiana per la Ricerca sul Cancro Foundation, Special Program Molecular Clinical Oncology, 5 1000, number 10007, Milan, Italy (to G.G. and to R.F.); Progetto Futuro in Ricerca 2008 (to D.R.); Programmi di Ricerca di Rilevante Interesse Nazionale (PRIN) 2008 (to G.G. and R.M.); PRIN 2009 (to D.R.); Progetto Futuro in Ricerca 2012 (to D.R.); Ministero dell'Istruzione, dell'Universita` e della Ricerca, Rome, Italy; Progetto Giovani Ricercatori 2008 (to D.R.); Progetto Giovani Ricercatori 2008 (to D.R.); Progetto Giovani Ricercatori 2010 (to D.R.); Ricerca Sanitaria Finalizzata 2008 (to G.G.); Ministero della Salute, Rome, Italy; Novara-AIL Onlus Foundation, Novara, Italy (to G.G. and D.R.); Compagnia di San Paolo, Turin, Italy (to R.F.); Helmut Horten Foundation and San Salvatore Foundation, Lugano, Switzerland (to F.B.); Nelia et Amadeo Barletta Foundation, Lausanne (to F.B.); National Institutes of Health grant PO1-CA092625 (to R.D.-F.); and a Specialized Center of Research grant from the Leukemia and Lymphoma Society (to R.D.-F.). S.M. and S.C. are supported by fellowships from the Novara-AIL Onlus Foundation, Novara, Italy. L.P. is on leave from the University of Perugia Medical School."
- "The authors declare no competing financial interests."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Yes	Appropriate data sources used: "multicentric cohort of 637 newly diagnosed and previously untreated CLL patients who consecutively presented for initial evaluation at four institutions from June 1996 through June 2011"
Domain 2: Predictors	Yes	Detailed description of predictors and timing of measurements: "clinical information prospectively collected at clinically relevant time points (i.e. at diagnosis, progression, and last follow-up)"
Domain 3: Outcome	Yes	Objective standard outcome; regular clinical database update
Domain 4: Analysis	Unclear	Unclear if individuals with missing data were excluded; number of events sufficient; building of model by recursive partitioning (decision tree, tenfold cross-validation), testing for stability with random forest algorithm and Cox model with bootstrapping; correction of c-statistic; assumptions were checked

Rossi V - Jeromin 2014 (Munich cohort)

Study characteristics

General information

Model and type of study

- Jeromin 2014 (MLL Munich cohort)
- Validation study

Secondary citations

· not applicable

Language of publication



Rossi V - Jeromin 2014	(Munich cohort	(Continued)
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• English

Study design

prospective cohort

Follow-up time

• median 55.2 months (range not reported)

Participants

Number of included participants in the cohort

• 1160 participants

Setting

· probably Germany, unclear

Recruitment period

• 2005 - 2010

Age (in years)

• median 67 years (range: 29.6 - 90.5 years)

Sex

• 64.6% male

Stages of disease

· not reported

Treatment

· Not reported

Inclusion criteria

· not reported

Exclusion criteria

· prior treatment

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- TP53 (binary); BIRC3 DIS (binary); SF3B1 M (binary); NOTCH1 M (binary); del11q22-q23 (binary)
- BIRC3, which is part of the original model, was not analysed in this study.

Timing of predictor measurement

• at diagnosis for most participants (82.6%)

Outcome(s)

Primary outcome in study

• overall survival: OS was measured from the date of diagnosis until last follow-up or death.

<u>Additional outcome(s)</u>

• time to treatment: TTT was evaluated from the date of diagnosis until the date of initial treatment.

Outcome in model development



Rossi V - Jeromin 2014 (Munich cohort) (Continued)

overall survival

Missing data

Participants with any missing data?

• yes (160 participants)

If yes, how was missing data handled?

• complete-case analysis

Analysis

Number of participants & number of events (specific time points where reported)

· overall survival: 930 participants, events not reported

Which model was used?

• original model as developed in Rossi 2013

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported (c-statistic)

Creation of risk groups?

- yes, 4 risk groups (cut-points not reported)
- at 5 years
- percentage of group without event at time point (low risk: 91%, intermediate risk: 90%, high risk: 75.2%, very high risk: 62.1%)

PROBAST: Applicability

Domain 1: Participant selection

• low

Domain 2: Predictors

low

Domain 3: Outcome

• low

Notes

Funding & conflict of interest

- "Next-generation deep sequencing studies were supported in part by the IRON-II study framework (Roche Diagnostics, Penzberg, Germany)."
- "SS, WK, CH, and TH are part owners of the MLL Munich Leukemia Laboratory. SJ, SW, VG, KB, FD, TA, AR and AK are employed by the MLL Munich Leukemia Laboratory."

Other comments

none

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Study design unclear



Rossi V - Jeromin 2014 (M	unich cohort) (Cor	ntinued)
Domain 2: Predictors	Yes	Relatively short recruitment period (2005 - 2010); samples were sent to a laboratory which was not involved in treatment.
Domain 3: Outcome	Yes	Objective standard outcome; no information on outcome assessment. However, we assumed that for OS, risk of bias was low.
Domain 4: Analysis	No	Individuals with missing data excluded from analysis (complete data for OS in 935/1160 cases); no performance measures, just risk groups
Overall judgement	No	

Rossi V - Rossi 2013 (unclear)

Stuay characteristics	study cha	racteristics
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General information	General	information
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Model and type of study

- Rossi 2013 (unclear)
- Validation study

Secondary citations

• not applicable

Language of publication

English

Study design

retrospective cohort

Follow-up time

• median 70.8 months, (range not reported)

Participants

Number of included participants in the cohort

· not reported

Setting

• Italy, multicentre study

Recruitment period

• 1996 - 2011

Age (in years)

not reported

Sex

• 55.9% male

Stages of disease

• Rai 0-I: 75.4%; Rai II: 17.3%; Rai III-IV: 7.3%

Treatment



Rossi V - Rossi 2013 (unclear) (Continued) • not reported
	Inclusion criteria
	not reported
	Exclusion criteria
	• not reported
Predictors	List of predictors used for validation (and changes between original predictors and predictors in validation study)
	 TP53 (binary); BIRC3 DIS (binary); SF3B1 M (binary); NOTCH1 M (binary); del11q22-q23 (binary) same predictors
	Timing of predictor measurement
	at diagnosis
Outcome(s)	Primary outcome in study
	overall survival
	Additional outcome(s)
	not applicable
	Outcome in model development
	overall survival
Missing data	Participants with any missing data?
	• unclear
	If yes, how was missing data handled?
	not applicable
Analysis	Number of participants & number of events (specific time points where reported)
	overall survival: 370 participants, 62 events
	Which model was used?
	recursive partitioning
	Was the model updated?
	yesWhat has been updated? Excel sheet
	Performance measures reported?
	Calibration: reported (observed-expected ratio)Discrimination: reported (c-statistic)
	Creation of risk groups?
	 yes, 4 risk groups, (cut-points at very low risk: del13q14 sole lesion; low risk: +12 or all lesions; intermediate: NOTCH1 and/or SF3B1; high risk: TP53 and/or BIRC3) at 5 years



Rossi V - Rossi 2013 (unclear) (Continued)

• percentage of group without event at time point (low risk: 86.9%, intermediate risk: 77.6%, high risk: 65.9%, very high risk: 50.9%)

PROBAST: Applicability

Domain 1: Participant selection

low

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "This study was supported by the Associazione Italiana per la Ricerca sul Cancro Foundation, Special Program Molecular Clinical Oncology, 5 1000, number 10007, Milan, Italy (to G.G. and to R.F.); Progetto Futuro in Ricerca 2008 (to D.R.); Programmi di Ricerca di Rilevante Interesse Nazionale (PRIN) 2008 (to G.G. and R.M.); PRIN 2009 (to D.R.); Progetto Futuro in Ricerca 2012 (to D.R.); Ministero dell'Istruzione, dell'Università e della Ricerca, Rome, Italy; Progetto Giovani Ricercatori 2008 (to D.R.); Progetto Giovani Ricercatori 2010 (to D.R.); Ricerca Sanitaria Finalizzata 2008 (to G.G.); Ministero della Salute, Rome, Italy; Novara-AIL Onlus Foundation, Novara, Italy (to G.G. and D.R.); Compagnia di San Paolo, Turin, Italy (to R.F.); Helmut Horten Foundation and San Salvatore Foundation, Lugano, Switzerland (to F.B.); Nelia et Amadeo Barletta Foundation, Lausanne (to F.B.); National Institutes of Health grant PO1-CA092625 (to R.D.-F.); and a Specialized Center of Research grant from the Leukemia and Lymphoma Society (to R.D.-F.). S.M. and S.C. are supported by fellowships from the Novara-AIL Onlus Foundation, Novara, Italy. L.P. is on leave from the University of Perugia Medical School."
- "The authors declare no competing financial interests."

Other comments

difference between external validation cohort and development cohort unclear

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	Unclear	Study design not clear: "The validation series was represented by a cohort of 370 newly diagnosed and previously untreated CLL patients who consecutively presented for initial evaluation from June 1996 through June 2011 (Table S1) and provided with regular follow-up (at least three visits/year)."
Domain 2: Predictors	Yes	No detailed description of predictor assessment; we assumed that it was similar to the development cohort described in the same publication.
Domain 3: Outcome	Yes	Objective standard outcome, regular clinical database update
Domain 4: Analysis	No	Small number of events; no information on missing values; calibration not reported
Overall judgement	No	

Stephens OS D - Stephens 2015 (Ohio cohort)

Study characteristics



Stephens OS D - Stephens 2015 (Ohio cohort) (Continued)

General information

Model and type of study

- Stephens 2015
- · Development study

Secondary citations

not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time

· not reported

Participants

Number of included participants in the cohort

• 114 participants

Setting

• USA, single-centre study (Ohio State University)

Recruitment period

• 2002 - 2012

Age (in years)

• median 62 years (range: 40 - 92 years)

Sex

· not reported

Stages of disease

• Rai 0: 33%; Rai I-II: 46%; Rai III-IV: 21%

<u>Treatment</u>

- 35 received purine analogue-based therapy
- 18 received treatment on a clinical trial
- 5 received other treatment

Inclusion criteria

• patients with CLL with del(17p13.1), previously untreated

Exclusion criteria

· not reported

Predictors

Number of candidate predictors

- univariable analysis: no univariable analysis
- multivariable analysis: 18

List of predictors in final model (including cut-points for dichotomised factors)



Stephens OS D - Stephens 2015 (Ohio cohort) (Continued)

• for OS: ECOG Performance Status (binary); age (binary); lactate dehydrogenase (LDH; binary)

Timing of predictor measurement

• at presentation, median 4.8 months after diagnosis (0 days to 19.7 years)

Outcome(s)

Primary outcome in study

overall survival: OS was calculated from date of first visit until date of death or last follow-up.

Additional outcome(s)

• treatment-free survival: TFS was calculated from date of first visit until date of first treatment or death, censoring patients alive and treatment-free at last follow-up.

Missing data

Participants with any missing data?

yes

If yes, how was missing data handled?

· multiple imputation technique

Analysis

Number of participants & number of events (specific time points where reported)

- · OS: 114 participants, events not reported
- TFS: 114 participants, 58 events

Predictor selection method

• univariable analysis, during multivariable modelling: backwards selection

Statistical method

· proportional hazards model

Simplification of model?

- yes
- risk score (RS) based on the variables and regression coefficients of the multivariable model
- simplified risk score (SRS) to be used in clinical practice based on the strength of associations with clinical outcome when all variables had been categorised

Performance measures reported?

- Calibration: not reported
- Discrimination: reported for OS (c-statistic), but not reported for TFS (c-statistic for multivariable model only)

Creation of risk groups?

- for OS: yes, 3 risk groups (SRS = 0; 2; 4)
- at 2 years

PROBAST: Applicability

Domain 1: Participant selection

• high - selective sample of very high risk individuals with a del(17p) deletion

<u>Domain 2: Predictors</u>

low

Domain 3: Outcome



Stephens OS D - Stephens 2015 (Ohio cohort) (Continued)

low

Notes

Funding & conflict of interest

- "Contract grant sponsors: Four Winds Foundation, D. Warren Brown Foundation; Contract grant sponsors: Mr. and Mrs. Michael Thomas, Harry Mangurian Foundation; Contract grant no.: P50 CA140158; Contract grant sponsor: Leukemia and Lymphoma Society".
- "The authors have no competing interests to disclose."

Other comments

The authors reported a multivariable model, a risk score (RS) and a simplified risk score (SRS) for each
outcome. Since the former two were not validated externally, only data for the SRS were extracted.

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Possibly selective inclusion of individuals, as only patients with del(17p13.1) were considered. Individuals without assessment could not have been included. Other eligibility criteria were unclear.
Domain 2: Predictors	Yes	Mostly laboratory data, probably assessed in a similar way: "Stimulated cytogenetic and fluorescent in situ hybridisation (FISH) analyses were performed on peripheral blood or bone marrow samples, as previously described"; single-centre study
Domain 3: Outcome	Yes	Objective standard outcome: although median observation time was not reported, median OS estimates were 5.2 years, therefore, we assumed a median observation time exceeding 5 years.
Domain 4: Analysis	No	Number of events only reported for TFS, not OS - the number of persons included was too small, total n = 114; missing data handling appropriate; inconsistent reporting of c-statistics; calibration was not reported
Overall judgement	No	

Stephens OS V - Stephens 2015 (MDACC)

Study characteristics

General	information

Model and type of study

- · Stephens 2015
- · Validation study

Secondary citations

not applicable

Language of publication

• English

Study design

· retrospective cohort

Follow-up time



Stephens OS V - Stephens 2015 (MDACC) (Continued)

· not reported

Participants

Number of included participants in the cohort

• 129 participants

Setting

• USA, single-centre study (MDACC)

Recruitment period

· not reported

Age (in years)

• median 63 years (range: 40 - 85 years)

Sex

not reported

Stages of disease

• Rai 0: 29%; Rai I-II: 52%; Rai III-IV: 19%

Treatment

· not reported

Inclusion criteria

• patients with CLL with del(17p13.1), previously untreated

Exclusion criteria

· not reported

Predictors

<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>

- for OS: ECOG Performance Status (binary); age (binary); lactate dehydrogenase (LDH; binary)
- · same predictors

Timing of predictor measurement

not reported

Outcome(s)

Primary outcome in study

overall survival

Additional outcome(s)

treatment-free survival

Outcome in model development

- overall survival
- treatment-free survival

Missing data

Participants with any missing data?

not reported

If yes, how was missing data handled?



Stephens OS V - Stephens 2015 (MDACC) (Continued)

· not applicable

Analysis

Number of participants & number of events (specific time points where reported)

- · OS: 129 participants, events not reported
- TFS: 129 participants, events not reported

Which model was used?

· simplified risk score (SRS)

Was the model updated?

no

Performance measures reported?

- Calibration: not reported
- Discrimination: reported for OS and TFS (c-statistic)

Creation of risk groups?

- for OS: yes, 3 risk groups (SRS = 0; 2; 3)
- at 2 years

PROBAST: Applicability

Domain 1: Participant selection

• high - selective sample of very high risk individuals with a del(17p) deletion

Domain 2: Predictors

low

Domain 3: Outcome

low

Notes

Funding & conflict of interest

- "Contract grant sponsors: Four Winds Foundation, D. Warren Brown Foundation; Contract grant sponsors: Mr. and Mrs. Michael Thomas, Harry Mangurian Foundation; Contract grant no.: P50 CA140158; Contract grant sponsor: Leukemia and Lymphoma Society".
- "The authors have no competing interests to disclose."

Other comments

none

Item	Authors' judgement Support for judgement	
Domain 1: Participant se- lection	No	Possibly selective inclusion of individuals, as only patients with del(17p13.1) were considered. Individuals without assessment could not have been included. Other eligibility criteria were unclear.
Domain 2: Predictors	Unclear	lack of information about similarity of predictor assessment to the development study; no recruitment period
Domain 3: Outcome	Yes	Objective standard outcome: although median observation time was not reported, median OS estimates were 6.4 years, therefore we assumed a median observation time exceeding 5 years.



Stephens	OSV-	Stephens	2015	(MDACC)	(Continued)
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Domain 4: Analysis	No	Number of included persons too small, total N = 129; calibration not reported
Overall judgement	No	

Stephens TFS D - Stephens 2015 (Ohio cohort)

Study characteristics	
General information	Model and type of study
	Stephens 2015
	Development study
	Different outcome but identical publication as Stephens OS D - Stephens 2015 (Ohio cohort) and Stephens OS V - Stephens 2015 (MDACC). Therefore, only relevant differences are reported in this table For additional information, see tables above.
Participants	see Stephens OS D - Stephens 2015 (Ohio cohort)
Predictors	List of predictors in final model (including cut-points for dichotomised factors)
	 for TFS: ECOG Performance Status (binary); Rai stage (binary); age (binary); white blood cell (WBG binary); del(11q22.3) (binary)
Outcome(s)	Primary outcome in study
	• overall survival: OS was calculated from date of first visit until date of death or last follow-up.
	Additional outcome(s)
	 treatment-free survival: TFS was calculated from date of first visit until date of first treatment or deatl censoring patients alive and treatment-free at last follow-up.
Missing data	
Analysis	Number of participants & number of events (specific time points where reported)
	OS: 114 participants, events not reported
	TFS: 114 participants, 58 events
	Performance measures reported?
	Calibration: not reported
	 Discrimination: reported for OS (c-statistic), but not reported for TFS (c-statistic for multivariable model only)
	Creation of risk groups?
	 for TFS: yes, 3 risk groups (SRS = 0/1; 2/3; >= 4) at 2 years
PROBAST: Applicability	
Notes	



Stephens TFS D - Stephens 2015 (Ohio cohort) (Continued)

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Possibly selective inclusion of individuals, as only patients with del(17p13.1) were considered. Individuals without assessment could not have been included. Other eligibility criteria were unclear.
Domain 2: Predictors	Yes	Mostly laboratory data, probably assessed in a similar way: "Stimulated cytogenetic and fluorescent in situ hybridization (FISH) analyses were performed on peripheral blood or bone marrow samples, as previously described"; single-centre study
Domain 3: Outcome	No	Predictors were not excluded from the outcome definition (Rai stage; Hallek 2008)
Domain 4: Analysis	No	Number of events only reported for TFS, not OS - the number of persons included was too small, total N = 114; missing data handling appropriate; inconsistent reporting of c-statistics; calibration was not reported
Overall judgement	No	

Stephens TFS V - Stephens 2015 (MDACC)

Study characteristics	
General information	Model and type of study
	Stephens 2015Validation study
	Different outcome but identical publication as Stephens OS D - Stephens 2015 (Ohio cohort) and Stephens OS V - Stephens 2015 (MDACC). Therefore, only relevant differences are reported in this table. For additional information, see tables above.
Participants	
Predictors	<u>List of predictors used for validation (and changes between original predictors and predictors in validation study)</u>
	 for TFS: ECOG Performance Status (binary); Rai stage (binary); age (binary); white blood cell (WBC; binary); del(11q22.3) (binary) same predictors
Outcome(s)	Primary outcome in study
	overall survival
	Additional outcome(s)
	treatment-free survival
	Outcome in model development
	overall survivaltreatment-free survival
Missing data	



Stephens TFS V - Stephens 2015 (MDACC) (Continued)

Analysis

Number of participants & number of events (specific time points where reported)

- OS: 129 participants, events not reported
- TFS: 129 participants, events not reported

Performance measures reported?

- · Calibration: not reported
- Discrimination: reported for OS and TFS (c-statistic)

Creation of risk groups?

- for TFS: yes, 3 risk groups (SRS = 0/1; 2/3; >= 4)
- at 2 years

PROBAST: Applicability

Notes

Item	Authors' judgement	Support for judgement
Domain 1: Participant se- lection	No	Possibly selective inclusion of individuals, as only patients with del(17p13.1) were considered. Individuals without assessment could not have been included. Other eligibility criteria were unclear.
Domain 2: Predictors	Unclear	Lack of information about similarity of predictor assessment to the development study, no recruitment period
Domain 3: Outcome	No	Predictors were not excluded from the outcome definition (Rai stage; Hallek 2008)
Domain 4: Analysis	No	Number of included persons too small, total N = 129; calibration not reported
Overall judgement	No	

AIHA: autoimmune haemolytic anaemia

ALC: absolute lymphocyte count

AUC: area under the curve B2M: beta2-microglobulin

 $BcR\ IG: clonotypic\ B-cell\ receptor\ immunoglobulin$

BIRC3: baculoviral IAP repeat containing 3

BR: bendamustine-rituximab C: cyclophosphamide

CHL: chlorambucil

CHOP: cyclophosphamide, doxorubicin, vincristine, and prednisone

CI: confidence interval

CIRS: Cumulative Illness Rating Score CLL: chronic lymphocytic leukaemia

CLL-IPI: Chronic Lymphocytic Leukaemia-International Prognostic Index

cMBL: clinical monoclonal B lymphocytosis

CVP: central venous pressure

DE: Germany

del11q: FISH-detected genetic abberation del11q del17p: FISH-detected genetic abberation del17p

ECOG PS: Eastern Cooperative Oncology Group performance status

F: fludarabine

FC: fludarabine and cyclophosphamide



FCR: fludarabine, cyclophosphamide and rituximab

FFP: fresh frozen plasma

FR: France

FISH: fluorescence in situ hybridisation HIV: human immunodeficiency virus

HBV: hepatitis B virus HCV: hepatitis C virus

HDMP: high-dose methylprednisolone

IgHV: immunoglobulin heavy chain variable region genes

IGLV: immunoglobulin lambda variable cluster

IQR: Interquartile range IPD: individual patient data

iwCLL: International workshop on Chronic Lymphocytic Leukemia

LDH: lactate dehydrogenase LDT: lymphocyte doubling time

LN: lymph node

LYM: lymphocyte count MCAR: missing completely at random

M-CLL: mutated chronic lymphocytic leukaemia

MDACC: MD Anderson Cancer Center MLL: Munich Leukemia Laboratory MoAbs: monoclonal antibodies MV analyses: multivariable analysis

NA: not available

NCI: National Cancer Institute NOTCH1: notch receptor 1

O-Benda: ofatumumab bendamustine

O:E: observed-expected ratio

OS: overall survival

PFS: progression-free survival

PL: Poland

PROBAST: Prediction model Risk Of Bias ASsessment Tool

R: rituximab

RCT: randomised controlled trial

ROC: receiver operating characteristics curve

RR: risk ratio RS: risk score

RTK: tyrosine kinase inhibitors s-B2M: serum beta-2 microglobulin

SCALE: Scandinavian population-based study

SCAN: Scandinavian population-based case-control study

SF3B1: Splicing Factor 3b Subunit 1

SRS: simplified risk score s-TK: serum thymidine kinase TFS: treatment-free survival TP53: tumour protein P53 TP: treatment probability TTE: Time-to-Evenr

TTFT: time-to-first-treatment

U-CLL: unmutated chronic lymphocytic leukaemia

WBC: white blood cell count WHO: World Health Organization

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Apelgren 2006	Study focus on staging system(s)
Baccarani 1982	Study focus on staging system(s)



Study	Reason for exclusion
Baliakas 2015	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Berke 2019	Prognostic factor study
Bettini 1986	Outdated staging system, which is no longer used in today's clinical practice
Binet 1977	Study focus on staging system(s)
Binet 1981	Study focus on staging system(s)
Bo 2014	Prognostic factor study
Bomben 2005	Comment or addition referring to an excluded study
Bomben 2009	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Bou Samra 2014	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Brejcha 2010	Prognostic factor study
Brugiatelli 2007	Prognostic factor study
Bulian 2014	Prognostic factor study
Byrd 2006	Prognostic factor study
Cailliod 2005	Prognostic factor study
Callea 1999	Prognostic factor study
Catovsky 1989	Prognostic factor study
Cesano 2013	Prognostic factor study
Chang 2003	Prognostic factor study
Chastang 1985	Outdated staging system, which is no longer used in today's clinical practice
Chauzeix 2018	Prognostic factor study
Chelazzi 1979	Study focus on staging system(s)
Chen 1997	Prognostic factor study
Chena 2008	Prognostic factor study
Chevallier 2002	Prognostic factor study
Chiaretti 2014	Prognostic factor study
Christiansen 1994	Prognostic factor study
Chuang 2012	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Ciccone 2012	Prognostic factor study



Study	Reason for exclusion
Ciocoiu 1988	Study focus on staging system(s)
Claus 2012	Prognostic factor study
Claus 2014	Prognostic factor study
Cmunt 2002	Prognostic factor study
Cocco 2005	Prognostic factor study
Corcoran 2005	Prognostic factor study
Cordone 1998	Prognostic factor study
Cortese 2014	Prognostic factor study
Coscia 2012	Prognostic factor study
Crespo 2003	Prognostic factor study
Criel 1997	Characterisation of CLL and prognostic factors
Cro 2009	Prognostic factor study
Cro 2010	Characterisation of CLL and prognostic factors
Cuneo 2004	Characterisation of CLL and prognostic factors
Cuneo 2018	Observational efficacy study
D'Arena 2001	Prognostic factor study
D'Arena 2007	Prognostic factor study
D'Arena 2012	Characterisation of CLL and prognostic factors
Damle 1999	Prognostic factor study
Dasgupta 2015	Study focus on identification of the threshold for a prognostic factor
Davis 2016	Study focus on identification of the threshold for a prognostic factor
De Faria 2000	Study focus on staging system(s)
De Rossi 1989	Study focus on staging system(s)
DeAndres-Galiana 2016	Study focus on identification of a prognostic factor
Degan 2004	Prognostic factor study
Del Guidice 2011	Prognostic factor study
Del Poeta 2010	Prognostic factor study
Del Principe 2004	Prognostic factor study



Study	Reason for exclusion
Del Principe 2006	Prognostic factor study
Delgado 2009	Prognostic factor study
Delgado 2014	Prognostic factor study
Deslandes 2007	Methodological study
Di Raimondo 2001	Prognostic factor study
Dimier 2018	Prediction model to evaluate the relationship between surrogacy outcome(s) and standard outcomes
Dong 2011	Prognostic factor study
Dong 2014	Prognostic factor study
Durak 2009	Prognostic factor study
El-Kinawy 2012	Prognostic factor study
Fang 2019	Study applies the CLL-IPI, but without the aim of validating it
Ferrara 1981	Study focus on staging system(s)
Ferreira 2014	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
French Cooperative Group on CLL 1988	Outdated staging system, which is no longer used in today's clinical practice
Friedman 2009	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Gattei 2008	Prognostic factor study
Gdynia 2018	Prognostic factor study
Geisler 1997	Characterisation of CLL and prognostic factors
Gentile 2016	Prognostic factor study
Gentile 2018	Score or model included predictors available only at treatment initiation
Giles 2003	Patient population did not match the review question; study included previously treated CLL patients
Giudice 2018	Prognostic factor study
Gogia 2014	Prognostic factor study
Gonzalez 2013	Characterisation of CLL and prognostic factors
Gonzalez-Gascon 2015	Characterisation of CLL and prognostic factors
Gonzalez-Rodriguez 2010	Characterisation of CLL and prognostic factors



Study	Reason for exclusion
Grabowski 2005	Prognostic factor study
Grever 2006	Abstract only
Hallek 1999	Characterisation of CLL and prognostic factors
Han 1984	Prognostic factor study
Herold 2011	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Hock 2010	Prognostic factor study
Houldsworth 2014	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Hus 2006	Prognostic factor study
Jaksic 1981	Prognostic factor study
Jaksic 1992	Study focus on staging system(s)
Jaksic 2014	Comment or addition referring to an excluded study
Josefsson 2007	Prognostic factor study
Juliusson 1986	Prognostic factor study
Juliusson 1990	Prognostic factor study
Kahraman 2014	Prognostic factor study
Kardum-Skelin 2008	Prognostic factor study
Kardum-Skelin 2009	Patient population did not match the review question; study included previously treated CLL patients
Karmiris 1994	Prognostic factor study
Kay 2018	Study applied the CLL-IPI, but without the aim of validating it
Keating 2000	Patient population did not match the review question; study included previously treated CLL patients
Khalifa 2002	Prognostic factor study
Kim 2004	Prognostic factor study
Kimby 1988	Prognostic factor study
Kleinstern 2018	Risk factor study
Knospe 1977	Prognostic factor study
Koberda 1989	Prognostic factor study
Korycka-Wolowiec 2011	Prognostic factor study



Study	Reason for exclusion
Krober 2002	Prognostic factor study
Krober 2006	Patient population did not match the review question; study included previously treated CLL patients
Kryachok 2011	Prognostic factor study
Kurec 1992	Prognostic factor study
Lai 2002	Prognostic factor study
Lech-Maranda 2012	Prognostic factor study
Lech-Maranda 2013	Prognostic factor study
Lecouvet 1997	Prognostic factor study
Li 2008	Prognostic factor study
Li 2017a	Prognostic factor study
Lin 2002	Prognostic factor study
Lin 2014	Prognostic factor study
Lozano-Santos 2014	Prognostic factor study
Lucas 2015	Prognostic factor study
Maffei 2007	Prognostic factor study
Maffei 2010	Prognostic factor study
Mandelli 1987	Development study of an outdated staging system, which is no longer used in today's clinical prac- tice
Mansouri 2013	Prognostic factor study
Marasca 2005	Prognostic factor study
Marasca 2013	Prognostic factor study
Martinelli 2008	Prognostic factor study
Masic 1998	Prognostic factor identification study
Mateva 2001	Prognostic factor model, no full-text for further investigation
Matthews 2006	Prognostic factor study
Matthews 2007	Prognostic factor study
Matutes 2013	Prognostic factor study
Matutes 2017	Review



Study	Reason for exclusion
Melo 1987	Patient population did not match the review question; study was not restricted to previously untreated CLL patients
Miao 2018	Prognostic factor study
Molica 1984	Study focus on staging system(s)
Molica 1986	Prognostic factor study
Molica 1988	Prognostic factor study
Molica 1991	Prognostic factor study
Molica 1994	Prognostic factor study
Molica 1998	Prognostic factor study
Molica 1999a	Prognostic factor study
Molica 1999b	Prognostic factor study
Molica 2008	Prognostic factor study
Montserrat 1991	Prognostic factor study
Morabito 2001	Prognostic factor study
Morabito 2015a	Prognostic factor study
Morabito 2015b	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Morabito 2018c	Study aimed to identify cut-off points for a specific prognostic factor
Moreno 2019	Study focus on staging system(s)
Morilla 2008	Prognostic factor study
Nabhan 2017	Score or model included predictors available only at treatment initiation
Nedeva 2018	Evaluation of an excluded model
Nenova 2000	Prognostic factor study
Nipp 2014	Prognostic factor study
Nola 2004	Patient population did not match the review question; study was not restricted to previously untreated CLL patients
Nowakowski 2009	Prognostic factor study
Nuckel 2006	Prognostic factor study
Nuckel 2009	Prognostic factor study



Study	Reason for exclusion
O'Brien 1993	Patient population did not match the review question; study was not restricted to previously untreated CLL patients
Ocana 2007	Prognostic factor study
Oliveira 2011	Prognostic factor study
Orgueira 2019	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Oscier 1990	Prognostic factor study
Paolino 1984	Prognostic factor study
Plesingerova 2017	Study focus on surrogacy outcome(s)
Prokocimer 1985	Prognostic factor study
Qin 2017	Prognostic factor study
Queiros 2015	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Rai 1975	Study focus on staging system(s)
Rai 1990	Study focus on staging system(s)
Raponi 2018	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Resegotti 1989	Prognostic factor study
Rissiek 2014	Prognostic factor study
Rodriguez 2007	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Ronchetti 2016	Prognostic factor study
Rossi 1986	Study focus on staging system(s)
Rossi 2010a	Study focus on identification of the threshold for a prognostic factor
Rossi 2010b	Patient population did not match the review question; study was not restricted to previously untreated CLL patients
Rossi 2011	Patient population did not match the review question; study was not restricted to previously untreated CLL patients
Rozman 1979	Study focus on staging system(s)
Salomon-Nguyen 1995	Diagnostic score
Santoro 1979	Study focus on staging system(s)
Sarmiento 2002	Prognostic factor study
Savvopoulos 2016	Methodological study



Study	Reason for exclusion
Scolozzi 1981	Study focus on staging system(s)
Shanafelt 2010	Prognostic factor study
Shanafelt 2017	Outcome not relevant for the review
Spacek 2009	Prognostic factor study
Stamatopoulos 2009	Prognostic score, however, this score was built with a proxy as outcome ('distinguished mutated and unmutated cases')
Stamatopoulos 2017	Prognostic factor study
Strefford 2015	Prognostic factor study
Szymczyk 2018	Prognostic factor study
Tallarico 2018	Outcome not relevant for the review
Tobin 2005a	Study focus on identification of the threshold for a prognostic factor
Tobin 2005b	Study focus on identification of the threshold for a prognostic factor
Vallat 2013	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Van Damme 2012	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering
Velardi 1980	Study focus on staging system(s)
Vojdeman 2017	Prognostic factor study
Vural 2014	Prognostic factor study
Weinberg 2007	Patient population did not match the review question; study was not restricted to previously untreated CLL patients
Weiss 2011	Prognostic factor study
Wierda 2003	Prognostic factor study
Winkler 2010	Prognostic factor study
Wu 2010	Study focus on staging system(s)
Zengin 1997	Study focus on staging system(s)
Zenz 2009	Prognostic factor study
Zucchetto 2006	Genetic analysis, e.g. genetic subgrouping, genetic signature(s) or genetic clustering

CLL: chronic lymphocytic leukaemia

CLL-IPI: chronic lymphocytic leukaemia International Prognostic Index



Characteristics of studies awaiting classification [ordered by study ID]

Antic 2011	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Saumann 2014	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Bulian 2011	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Bulian 2012	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Cavallini 2017	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Del Guidice 2005	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Friedrichs 2011	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Furundarena 1994	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.



Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix
	7.
laferlach 2010	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Jarque 1991	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Lee 1987	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Leotard 2000	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Letestu 2010	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
.i 2017b	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Liang 2018	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.



Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix
Notes	7.
Miao 2019	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Molica 1990	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Molica 2010	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Molica 2015	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Molica 2019	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Morabito 2011	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Pepper 2012	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.



Qin 2018	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Rozman 1982	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Rozman 1984	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Schweighofer 2011	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Stamatopoulos 2010	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Tsimberidou 2007	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Vetro 2018	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.
Visentin 2015	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.



Wierda 2009	
Notes	Details on study design, recruitment period, prognostic factors and more can be found in Appendix 7.

Characteristics of ongoing studies [ordered by study ID]

NCT00275054

Study name	Rituximab, fludarabine, and cyclophosphamide or observation alone in treating patients with stage 0, stage I, or stage II chronic lymphocytic leukemia
Starting date	2005
Contact information	Michael Hallek, MD, Medizinische Universitaetsklinik I at the University of Cologne
Notes	 inclusion reason: "Development of a new prognostic staging system" no staging system or prognostic model published yet

NCT03436524

Study name A prognostic tool for early stage CLL					
Starting date	2018				
Contact information	Davide Rossi, MD, PhD, Principal investigator, Oncology Institute of Southern Switzerland				
Notes	 inclusion reason: "The study aims at developing a model for the prediction of time to first treatment in chronic lymphocytic leukemia patients presenting with asymptomatic early stage disease". study is currently recruiting 				

APPENDICES

Appendix 1. MEDLINE (via Ovid) search strategy up to 19 September 2018

1	exp LEUKEMIA, LYMPHOCYTIC, CHRONIC, B-CELL/
2	((lymphocytic* or b-cell or b-lymphocytic or lymphoblastic or lymphatic) adj1 leuk?em\$ adj3 (chronic\$ or cronic\$ or chroniq\$ or well-differentia\$)).tw,kf,ot.
3	((lymphoplasmacytoid or lymphocytic) adj1 lymphom* adj3 (chronic\$ or cronic\$ or chroniq\$ or well-differentia\$)).tw,kf,ot.
4	((small cell\$ or small-cell\$) adj3 lymphom\$).tw,kf,ot.
5	(lymphom\$ adj2 lymphocyt\$).tw,kf,ot.
6	lymphoplasma?ytoid.tw,kf,ot.



(Continued)								
7	(cll or b-cll or bcll).tw.							
8	sll.tw.							
9	or/1-8							
10	(predict\$ or clinical\$ or outcome\$ or risk\$).mp.							
11	validat\$.mp. or predict\$.ti. or rule\$.mp.							
12	(predict\$ and (outcome\$ or risk\$ or models\$)).mp.							
13	((history or variable\$ or criteria or scor\$ or characteristic\$ or finding\$ or factor\$) and (predict\$ or model\$ or decision\$ or identif\$ or prognos\$)).mp.							
14	decision\$.mp. and ((model\$ or clinical\$).mp. or LOGISTIC MODELS/)							
15	(prognostic and (history or variable\$ or criteria\$ or scor\$ or characteristic\$ or finding\$ or factor\$ or model\$)).mp.							
16	or/10-15							
17	validat\$.mp. or predict\$.ti. or DECISION SUPPORT TECHNIQUES/ or rule\$.mp. or PREDICTIVE VAL- UE OF TESTS/							
18	(predict\$ and (clinical\$ or identif\$)).mp.							
19	17 or 18							
20	RISK ASSESSMENT/							
21	(risk\$ adj scores\$).tw,kf,ot.							
22	exp RISK FACTORS/							
23	(risk\$ adj (score\$ and factor\$)).tw,kf,ot.							
24	DECISION SUPPORT TECHNIQUES/							
25	(decision\$ adj2 (techniqu\$ or model\$)).tw,kf,ot.							
26	(decision\$ and support\$ and technique\$).tw,kf,ot.							
27	(prediction\$ and rule\$ and clinical\$).tw,kf,ot.							
28	(decision\$ adj2 (modeling\$ or aid\$ or analys\$ or technique\$)).tw,kf,ot.							
29	or/20-28							
30	ANIMALS/ not HUMANS/							
31	9 and (16 or 19 or 29)							
32	31 not 30							



Appendix 2. MEDLINE (via Ovid) search strategy, 19 September 2018 to 24 June 2019, including study filter

1	exp LEUKEMIA, LYMPHOCYTIC, CHRONIC, B-CELL/
2	((lymphocytic* or b-cell or b-lymphocytic or lymphoblastic or lymphatic) adj1 leuk?em\$ adj3 (chronic\$ or cronic\$ or chroniq\$ or well-differentia\$)).tw,kf,ot.
3	((lymphoplasmacytoid or lymphocytic) adj1 lymphom* adj3 (chronic\$ or cronic\$ or chroniq\$ or well-differentia\$)).tw,kf,ot.
4	((small cell\$ or small-cell\$) adj3 lymphom\$).tw,kf,ot.
5	(lymphom\$ adj2 lymphocyt\$).tw,kf,ot.
6	lymphoplasma?ytoid.tw,kf,ot.
7	(cll or b-cll or bcll).tw.
8	sll.tw.
9	or/1-8
10	Validat\$.tw. or Predict\$.ti. or Rule\$.tw. or (Predict\$ and (Outcome\$ or Risk\$ or Model\$)).tw. or ((History or Variable\$ or Criteria or Scor\$ or Characteristic\$ or Finding\$ or Factor\$) and (Predict\$ or Model\$ or Decision\$ or Identif\$ or Prognos\$)).tw. or (Decision\$.tw. and ((Model\$ or Clinical\$).tw. or LOGISTIC MODELS/)) or (Prognostic and (History or Variable\$ or Criteria or Scor\$ or Characteristic\$ or Finding\$ or Factor\$ or Model\$)).tw. or ("Stratification" or "Discrimination" or "Discriminate" or c-statistic or "Area under the curve" or "AUC" or "Calibration" or "Indices" or "Algorithm" or "Multivariable").tw.
11	9 and 10
12	exp ANIMALS/ not HUMANS/
13	11 not 12
14	limit 13 to ed=20180919-20190624

key: exp # /: explode # MeSH subject heading, tw: text word, kf: keyword heading word, ot: original title, * or \$: truncation, ?: wildcard, adj#: adjacent within # number of words

search line #10: Geersing G-J, Bouwmeester W, Zuithoff P, Spijker R, Leeflang M, Moons KG, et al. (2012) Search filters for finding prognostic and diagnostic prediction studies in Medline to enhance systematic reviews. PLOS One 2012; 7(2): e32844, doi:10.1371/journal.pone.0032844

Appendix 3. Embase search strategy up to 24 June 2019

1 exp CHRONIC LYMPHATIC LEUKEMIA/



(Continued)	
2	((lymphocytic* or b-cell or b-lymphocytic or lymphoblastic or lymphatic) adj1 leuk?em* adj3 (chronic* or cronic* or chroniq* or well-differentia*)).tw,kw.
3	((lymphoplasmacytoid or lymphocytic) adj1 lymphom* adj3 (chronic* or cronic* or chroniq* or well-differentia*)).tw,kw.
4	((small cell* or small-cell*) adj3 lymphom*).tw,kw.
5	(lymphoma* adj2 lymphocyt*).tw,kw.
6	lymphoplasma?ytoid.tw,kw.
7	(cll or b-cll or bcll).tw.
8	sll.tw.
9	or/1-8
10	Validat*.tw. or Predict*.ti. or Rule*.tw. or (Predict* and (Outcome* or Risk* or Model*)).tw. or ((History or Variable* or Criteria or Scor* or Characteristic* or Finding* or Factor*) and (Predict* or Model* or Decision* or Identif* or Prognos*)).tw. or (Decision*.tw. and ((Model* or Clinical*).tw. or STATISTICAL MODEL/)) or (Prognostic and (History or Variable* or Criteria or Scor* or Characteristic* or Finding* or Factor* or Model*)).tw. or ("Stratification" or "Discriminate" or c-statistic or "Area under the curve" or "AUC" or "Calibration" or "Indices" or "Algorithm" or "Multivariable").tw.
11	9 and 10
12	exp ANIMAL/ not HUMAN/
13	11 not 12
14	limit 13 to embase
15	limit 14 to em=197401-201926

key: exp # /: explode # MeSH subject heading, tw: text word, kw: keyword, *: truncation, ?: wildcard, adj#: adjacent within # number of words searchline #10: Geersing G-J, Bouwmeester W, Zuithoff P, Spijker R, Leeflang M, Moons KG, et al. Search filters for finding prognostic and diagnostic prediction studies in Medline to enhance systematic reviews. PLOS One 2012; 7(2): e32844, doi:10.1371/journal.pone.0032844 (translated in Embase)

Appendix 4. ClinicalTrials.gov search strategy up to 05 March 2020

Basic search

prognostic OR predictive OR model OR score | chronic lymphocytic leukemia OR CLL

Advanced search

 $Intervention/treatment: prognostic \ {\tt OR} \ predictive \ {\tt OR} \ {\tt Model} \ {\tt OR} \ score$

Recruitment: all studies
Study type: all studies

Appendix 5. WHO ICTRP search strategy

Basic search

Chronic lymphocytic leukemia AND prognostic



Chronic lymphocytic leukemia AND predictive

Chronic lymphocytic leukemia AND model

Chronic lymphocytic leukemia AND score

CLL AND prognostic

CLL AND predictive

CLL AND model

CLL AND score

Advanced search

Condition: chronic lymphocytic leukemia OR CLL

Intervention: prognostic OR predictive Or model OR score

Recruitment status: ALL

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Author, year	Model (= 1) or Score (= 2)	Outcome	Recruit- ment peri- od	Number of patients	Study design	Number of predictors included in fi- nal model	Discrimina- tion	Calibration
Delgado 2017	2	OS	NR	524	Retrospective cohort	IgHV; FISH abnormalities (2)	0.682	Calibration plot for sur- vival at 5 years
International CLL IPI work- ing group	2	OS	1997 to 2007	3472	RCTs	IgHV; B2M; age; stage; TP53 (5)	0.723	NR
Morabito 2009	2	OS	NR	262	Unclear	IgHV; ZAP-70; CD38 (3)	NR	NR
Pflug 2014	2	OS	1997 to 2006	1223	RCTs	IgHV; B2M; age; gender; del17p; ECOG; sTK; del11q (8)	0.75	NR
Wierda 2007	1	OS	1981 to 2004	1674	Retrospective cohort	IgHV; B2M; age; stage; gender; ALC	0.84	Calibration plot for sur- vival at 5 years
Stephens 2015	2	OS	2002 to 2012	114	Retrospective cohort	age; LDH; ECOG (3)	0.73	NR
Molica 2005	2	PFS	1991 to 2000	1138	Retrospective cohort	stage; gender; lymphocytosis; LDT (4)	NR	NR
Baliakas 2019	2	TFS	NR	1900	Retrospective cohort	IgHV; TP53; FISH abnormalities; gender (4)	0.745	NR
Gentile 2016	2	TFS	2007 to NR	480	Prospective co- hort	IgHV; B2M; stage; ALC (4)	0.75	Hosmer-May test

(Continued) Rossi 2013	2	TFS	NR	673	Retrospective cohort	TP53; FISH abnormalities; SF3B1 (3)	0.642	Bias-correct- ed calibration slope, 0.965
Wierda 2011	1	TFS	2004 to 2009	930	Retrospective cohort	IgHV; LDH; number of involved lymph nodes; FISH abnormalities; lymph node size in neck; interaction term (6)	NR	NR
Stephens 2015	2	TFS	2002 to 2012	114	Retrospective cohort	age; stage; ECOG; del11q; WBC count (5)	0.84	NR

Abbreviations: OS: overall survival, PFS: progression-free survival; TFS: treatment-free survival; NR: not reported; RCT: randomised controlled trial; IgHV: immunoglobulin heavy chain variable region gene mutational status; B2M: beta 2 microglobulin; FISH: fluorescence in situ hybridisation genomic aberrations; Zap-70: zeta-chain-associated protein kinase 70; CD38: cluster of differentiation 38; LDH: lactate dehydrogenase; LPL: lipoprotein lipase; ECOG: Eastern Cooperative Oncology Group performance status

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Author, year	Model (=1) or Score (=2)	Outcome	Recruit- ment peri- od	Number of patients	Study design	Predictors included in final model (number)	Discrimina- tion	Calibration	
Baumann 2014	2	OS	1990 to 2012	949	Retrospective cohort	B2M; stage; Zap-70; Comorbirities	NR	NR	
						(4)			
Bulian 2012	1	OS	1996 to 2008	620	Retrospective	IgHV; B2M; age	0.79	NR	
					cohort	stage; FISH; gender			
						(6)			
Bulian 2011	1	OS	1996 to NR	1480	Retrospective	B2M; age; stage; gender	0.78	Calibration	
					cohort	(4)		plot for sur- vival at 5 years	
Furundarena	1	os	1973 to 1992	150	Retrospective cohort	age; stage; gender; splenomegaly	NR	NR	
1994						(4)			
Haferlach 2010	2	OS	2005 to 2008	399	NR	IgHV; age; TP53; translocation IGH@ on 14q32; number of chromosome aberrations based on CBA; WBC count	NR	NR	
						(6)			
Jarque 1991,	1	OS	1969 to 1988	187	Retrospective	Age; Spinal infiltration;	NR	NR	
model A					cohort	BUN			
						(3)			
Jarque 1991, model B	1	OS	1969 to 1988	187	Retrospective	Albumin;	NR	NR	
model B					cohort	Spinal infiltration;			
						BUN;			
						cervical adenopathies			
						(4)			

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Continued)								
Lee 1987	1, 2	OS	1970 to 1983	325	Retrospective cohort	Age; LDH; uric acid; alkaline phos- phatase; external lymphadenopathy	NR	NR
						(5)		
Liang 2018, model 3a	2	OS	2000 to 2014	501	Retrospective cohort	IgHV; B2M; stage; TP53; albumin; HBV infection	time-depen- dent ROC	NR
						(6)	curve	
Molica 1990	1	OS	NR	221	Unclear	age; stage; LAR	NR	O/E ratios
						(3)		
Molica 2019	1, 2	os	2000 to NR	108	Retrospective	age; ADL; CIRS	c = 0.70	NR
					cohort	(3)	(95% CI 0.53 to 0.87)	
Pepper 2012	1	OS	NR	1154	Unclear	IgHV; age; CD38; LDT	NR	NR
			(4)		(4)			
Rozman 1982	1	OS	NR	150	Unclear	Splenomegaly; lymphocytosis; anaemia; thrombocytopenia	NR	O/E proba bilities
						(4)		
Rozman 1984	1	OS	NR	329	Unclear	lymphadenopathy;	NR	O/E ratios
						haemoglobin; bone marrow pattern; hepatomegaly		
						(4)		
Stamatopou-	2	OS	NR	170	Unclear	ZAP-70; LPL; miR-29	NR	NR
los 2010						(3)		
Tsimberidou	2	OS	1985 to 2005	1893	Retrospective	Age; B2M; Del17p; albumin; creatinine	NR	NR
2007					cohort	(5)		
Visentin 2015	2	OS	1983 to 2013	608	Retrospective	IgHV; CD38; FISH abnormalities	0.88	NR
					cohort	(3)		

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(Continued)								
Wierda 2009	1, 2	OS	1985 to 2004	595	Unclear	Age; B2M; treatment (3)	NR	Concordance index for calibration curve (nomogram): 0.81
Friedrichs 2011	2	PFS	NR	134	NR	IgHV; CD38;	NR	NR
						iLR		
						(3)		
Leotard 2000	1	PFS	1985 to 1997	88	Prospective co-	B2M; LDH; albumin; sCD23	NR	NR
					hort	(4)		
Letestu 2010	1, 2	PFS	NR	339	NR	B2M; CD38; lymphocytosis; sTK	NR	NR
						(4)		
Antic 2011	2	TFS	NR	33	Unclear	B2M; LPL; sVEGF	NR	NR
						(3)		
Cavallini 2017	2	TFS	NR	125	NR	SF3B1; ERK1/2 phosphorylation	NR	NR
						(2)		
Del Guidice	2	TFS	2003 to 2004	201	Prospective co-	ZAP-70; CD38	NR	NR
2005					hort	(2)		
Gentile 2009	2	TFS	NR	222	NR	IgHV; B2M; CD38	NR	NR
						(3)		
Haferlach 2010	2	ТТТ	2005 to 2008	399	NR	IgHV; ATM deletion; translocation involving IGH@ locus on 14q32; number of chromosome abnormalities	NR	NR
						(4)		
Li 2017b	2	TFS	2007 to 2015	406	NR	IgHV; stage; Del17p	NR	NR
						(3)		

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(Continued)								
Liang 2018 model 3b	2	TFS	2000 to 2014	501	Retrospective cohort	IgHV; LD; lymphocytosis; platelets; HBV	time-depen- dent ROC curve	NR
						(5)	curve	
Metze 2000	2	TFS	1995 to 1998	57	Prospective co- hort	total tumor mass score; percentage of cells with 1 AgNOR cluster	NR	NR
						(2)		
Miao 2019,	1, 2	TFS	2000 to 2017	399	Retrospective	IgHV; B2M; Age; stage; TP53; SF3B1	AUC = 0.762	calibration
model A (CLL- IPI-S)					cohort	(6)		plot
Miao 2019,	1, 2	TFS	2000 to 2017	399	Retrospective	IgHV; B2M; stage; TP53; SF3B1	AUC = 0.773	calibration
model B (CLL- PI)					cohort	(5)		plot
Molica 2010	2	TFS	1998 to 2008	150	Retrospective cohort	IgHV; BAFF	0.86	NR
						(2)		
Molica 2015	1	TFS	NR	322	Unclear	IgHV; B-cell count; interaction term	NR	NR
						(3)		
Morabito 2011	2	TFS	NR	449	Retrospective cohort	stage; ZAP-70; FISH abnormalities; SFLC (kappa + delta)	NR	NR
						(4)		
Pepper 2012	1	TFS	NR	1154	Unclear	IgHV; age; LDT; CD38	NR	NR
						(4)		
Qin 2018	2	TTT	2008 to 2016	334	Retrospective cohort	IgHV; EBV DNA positive; stage; TP53; ALC; HBsAg+;	AUC = 0.768	NR
						(6)		
Schweighofer	1	TFS	NR	131	Retrospective	SKI (gene); SLAMF1 (gene)	NR	NR
2011					cohort	(2)		

(Continued)									4
Stamatopou-	2	TFS	NR	170	Unclear	ZAP-70; LPL; miR-29c	NR	NR	•
los 2010						(3)			
Vetro 2018	1	ТТТ	NR	171	Unclear	IgHV; SF3B1; percentage of B cells; genomic aberrations	NR	NR	_ 3
						(4)			
Wierda 2009	1	TFS	1985 to 2004	595	Unclear	Age; B2M; treatment; bone marrow lymphocytes	NR	NR	

(4)

Abbreviations: OS: overall survival, PFS: progression-free survival; TFS: treatment-free survival; TTT: time to treatment; NR: not reported; RCT: randomised controlled trial; IgHV: immunoglobulin heavy chain variable region gene mutational status; B2M: beta 2 microglobulin; FISH: fluorescence in situ hybridisation genomic aberrations; Zap-70: zeta-chain-associated protein kinase 70; CD38: cluster of differentiation 38; LDH: lactate dehydrogenase; LPL: lipoprotein lipase; ECOG: Eastern Cooperative Oncology Group performance status; LAR: lymphocyte accumulation time; sVEGF: soluble vascular endothelial growth factor; iLR: immature laminin receptor; CIRS: Cumulative Illness Rating Score; ADL: Katz Activity of Daily Living; EBV: Epstein-Barr virus; HBsAg: surface antigen of hepatitis B virus

Appendix 8. Reporting deficiencies in all included studies with at least one external validation study (development + validation studies, listed per model)

Prognostic model	Number of cohorts	Recruit- ment period reported in:	Study design reported in:	Observation time (range) reported in:	Total sam- ple size re- ported in:	Number of events re- ported in:	Calibration reported in:	Discrimination (95% CI or SE) reported in:
Baliakas (Baliakas 2019)	2	0	2	1 (1)	2	0	0	1 (1)
Barcelona-Brno (Delga	6	3	4	6 (6)	6	5	1	5 (3)
do 2017)								
CLL-IPI (Bahlo 2016)	11	9	9	10 (6)	10	9	5	8 (6)
GCLLSG (Pflug 2014)	2	1	2	2 (0)	2	2	2	2 (2)
GIMEMA (Molica 2005)	2	2	2	1 (1)	2	1	0	1 (1)
MDACC 2007 (Wierda 2007)	10	7	8	8 (9)	9	9	2	8 (7)
MDACC 2011 (Wierda 2011)	2	1	2	2 (2)	2	2	0	1 (1)
Morabito model (Morabito, 2009)	2	2	1	2 (2)	2	2	0	0 (0)
O-CLL1 model (Gentile 2016)	3	1	1	3 (2)	3	3	0	3 (2)
Rossi (Rossi 2013)	3	3	3	3 (0)	3	2	0	2 (0)
Stephens (Stephens 2015)	2	1	1	0 (0)	2	0	0	2 (0)
Stephens (Stephens 2015)	2	1	1	0 (0)	2	0	0	2 (0)



HISTORY

Protocol first published: Issue 1, 2016 Review first published: Issue 7, 2020

CONTRIBUTIONS OF AUTHORS

Nina Kreuzberger: screening and selection of studies, development of data extraction form and data extraction, characteristics of studies, 'Risk of bias' assessment, statistical analysis, writing and drafting of the review, communication with and between authors

Johanna AAG Damen: statistical analysis

Marialena Trivella: statistical input, screening and selection of studies, data extraction, 'Risk of bias' assessment

Lise J Estcourt: medical and content input, screening and selection of studies, data extraction, 'Risk of bias' assessment

Angela Aldin: screening and selection of studies, data extraction, 'Risk of bias' assessment

Lisa Umlauff: data extraction, 'Risk of bias' assessment, characteristics of included studies

Maria Vazquez Montes: prototype of data extraction form, support in data extraction, 'Risk of bias' assessment

Robert Wolff: 'Risk of bias' input

Karel GM Moons: methodological input on reviews of prognosis studies

Ina Monsef: search strategy development

Farid Foroutan: 'Risk of bias' input

Karl-Anton Kreuzer: medical and content input

Nicole Skoetz: protocol development, screening and selection of studies, data extraction, 'Risk of bias' assessment, extensive proofread

and comments on the review draft

DECLARATIONS OF INTEREST

Nina Kreuzberger: My institution received a grant from the Federal Ministry of Education and Research, Germany to conduct this review.

Johanna AAG Damen: none known

Marialena Trivella: I am working as a statistical editor in a number Cochrane groups, and I declare that my work as a statistical editor is independent to this published work where I participate. The University of Oxford received a small part of the grant from the Federal Ministry of Education and Research, Germany, as reimbursement for my time spent on the project.

Lise J Estcourt: My institution received a grant from the Federal Ministry of Education and Research, Germany to conduct this review.

Angela Aldin: My institution received a grant from the Federal Ministry of Education and Research, Germany to conduct this review.

Lisa Umlauff: My institution received a grant from the Federal Ministry of Education and Research, Germany to conduct this review.

Maria Vazquez Montes: I am sponsored by the BHF to contribute on a similar prognostic models review for heart failure which allowed me to effectively participate in the work under consideration.

Robert Wolff: As employee of Kleijnen Systematic Reviews, I was the lead author of a systematic review on prostate cancer, commissioned by Elekta, Nucletron.

Karel Moons: none known

Ina Monsef: none known

Farid Foroutan: none known

Karl-Anton Kreuzer: board member and consultant for AbbVie, Alexion, Amgen, Ariad, Baxter, Bayer Health Care, Biotest, Boehringer Ingelheim, Bristol Myers Squibb, Celgene, Chugai, Gilead, Glaxo-SmithKline, Grifols, Hexal, Janssen, Jazz Pharmaceuticals, Leo, Mundipharma, MSD, Novartis, Pfizer, Roche, Shire, Teva. Grants, fees, honoraria and travel grants from AbbVie, Alexion, Amgen, Ariad,



Baxter, Bayer Health Care, Biotest, Boehringer Ingelheim, Bristol Myers Squibb, Celgene, Chugai, Gilead, Glaxo-SmithKline, Grifols, Hexal, Janssen, Jazz Pharmaceuticals, Leo, Mundipharma, MSD, Novartis, Pfizer, Roche, Shire, Teva.

Nicole Skoetz: My institution received a grant from the Federal Ministry of Education and Research, Germany to conduct this review.

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Internal sources

- University Hospital of Cologne, Department I of Internal Medicine, Germany
- NHS Blood and Transplant, UK

External sources

· Grant by the Federal Ministry of Education and Research (Grant no. 01KG1711), Germany

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We changed the title of the review to clarify the included population (newly-diagnosed adults with CLL). In the protocol, the title was 'Prognostic models for chronic lymphocytic leukaemia: an exemplar systematic review and meta-analysis'.

Structure of the review

As prognosis reviews are a new review type within Cochrane, little guidance had been published at protocol stage. Recently, a review template has been developed and published by the Cochrane Prognosis Methods group, which we have adopted for this review (methods.cochrane.org/prognosis/our-publications). In addition, we have added two sections ('Selection of studies' and ''Risk of bias' assessment') to report our group decisions transparently.

Search methods

We did not search PubMed, because content from PubMed can be identified via MEDLINE.

We added a search for Embase based on editorial comments.

We added a search of the following databases for ongoing trials instead of the metaRegister of Controlled Trials:

- ClinicalTrials.gov
- World Health Organization International Clinical Trials Registry Platform (WHO ICTRP)

We did not search conference proceedings for abstracts, because the paucity of information would not allow us to apply the 'Risk of bias' tool. To ensure that we identified all validation studies corresponding to a developed model, we screened the citations of all included studies (Web of Science, October 2018).

Inclusion and exclusion of studies

Originally, we planned to include only models published after 1990. However, prognostic model studies are often conducted based on retrospective data, which sometimes include the analysis of blood samples frozen at diagnosis and analysed many years after. To give the same chance to studies with samples that were analysed immediately and after several years for inclusion in this review, we decided not to limit our search strategy based on date of publication. As a clarification, any studies that explicitly aimed at defining a new staging system was excluded during title and abstract screening.

We decided to include prognostic models with the outcomes 'time-to-first treatment' and 'treatment-free survival' as additional outcomes, because individuals with low-risk CLL can often live a long time without disease progression and the need for treatment. Usually, at disease progression, treatment is indicated, which means that these two outcomes are quite similar and both are meaningful for the patient.

After several rounds of discussion, we decided that genetic signatures are beyond the scope of this review because: a) we prespecified that we would exclude factor identification studies, which most of the genetic signature studies are to start with; b) they are not yet ready to be applied in clinical practice due to the unavailability of detailed genetic information that these signatures use and the complexity of the algorithms; and c) more often than not, it is not feasible to assess a genetic signature study in the same context as a prognostic model.

Analysis and 'Risk of bias' assessment

Instead of the CHARMS checklist, we used the recently published PROBAST-tool to assess the risk of bias of individual studies (Wolff 2019). The checklist does not originally aim at the assessment of risk of bias, but as a guide for critical appraisal and data extraction. However, at the protocol stage, no tool specific to 'Risk of bias' assessment for prognostic models had been published.



The analysis was not specified precisely in the protocol, as methods for systematic reviews of prognosis are just developing. As described in the protocol, we decided to follow the current recommendations of the Cochrane Prognosis Methods Group. We added a more detailed description in the section Data collection and analysis. In short, we meta-analysed the performance measures of the various external validation studies per model where data were available. We planned to summarise the measures of calibration. Unfortunately, calibration in the form of O:E ratios or calibration plots was rarely reported. Instead, many studies only presented the observed outcome frequency per subgroup at a specific time point. To gain an overview of this information among several external validations, during the review process and in collaboration with the Cochrane Prognososis Methods Group, we decided to represent the outcome frequency graphically together with the pooled outcome frequency in a table.

As we only meta-analysed the external validation studies of a prognostic model, we analysed only models that were externally validated several times. Although we have included all prognostic models and scores that we identified, we did not describe models without any external validation studies in detail, because it is not recommended to use prognostic scores or models without any testing in independent cohorts, especially when the development sample was small (Moons 2015; Steyerberg 2013). References to these studies can be found in the list of studies awaiting classification; further information can be found in Appendix 7.

In the protocol, we did not specify any sensitivity analyses. We decided post hoc to explore the effect of including area under the curve (AUC) as a performance measure for discrimination instead of the c-statistic (Figure 23), the effect of the Newcombe estimation method for 95% CIs of the c-statistic (Figure 24; Figure 25), and the difference between studies using the original predictor or a proxy (del(17p) instead of TP53 mutation (Figure 26).

GRADE

We did not apply GRADE as no GRADE guidance has yet been developed to assess certainty of evidence from meta-analysis of prognostic models.

NOTES

Parts of this review, especially the methods, are from the Cochrane Haematology standard template.

INDEX TERMS

Medical Subject Headings (MeSH)

Age Factors; Bias; Biomarkers, Tumor; Calibration; Confidence Intervals; Discriminant Analysis; Disease-Free Survival; Genes, p53 [genetics]; Immunoglobulin Heavy Chains [genetics]; Immunoglobulin Variable Region [genetics]; Leukemia, Lymphocytic, Chronic, B-Cell [*mortality] [pathology]; *Models, Theoretical; Neoplasm Staging; Prognosis; Progression-Free Survival; Receptors, Antigen, B-Cell [genetics]; Reproducibility of Results; Tumor Suppressor Protein p53 [genetics]

MeSH check words

Adult; Female; Humans; Male