

Figure S1. Missingness analysis of BMI with respect to age. The marginplot method in the VIM R package was used to analyze the distribution of BMI for patients with missing or non-missing age data, and the distribution of age for patients with missing or non-missing BMI. The X-axis represents age in years, and the Y axis represents BMI. A box plot summarizing BMI in patients for whom age data is missing (red) or present is shown on the y-axis. A box plot summarizing age of patients for whom BMI is missing (red) or present (blue) is shown on the x-axis. The distribution of age data for patients with missing BMI was similar to that for patients with non-missing BMI. Likewise, the distribution of BMI data for patients with missing age was similar to that for patients with non-missing age. The plot suggests BMI and age data are missing completely at random.

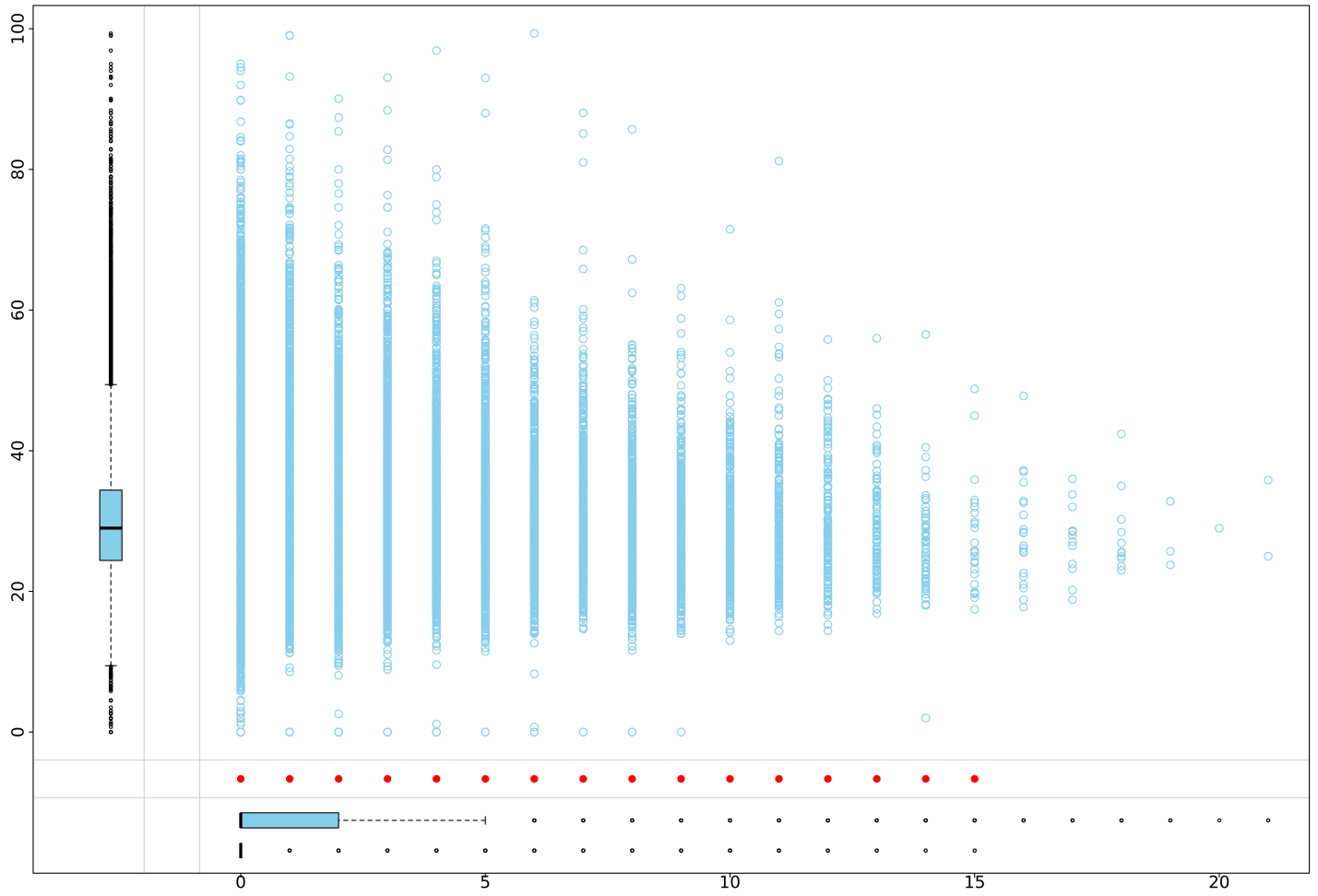


Figure S2. Missingness analysis of BMI with respect to Charleson severity index score. See Supplemental Figure S1 for explanations. The X axis represents the Charleson severity index score and the Y axis represents BMI.

COVID-19 severity	Percent of patients taking NSAIDs
Mild	3.6%
Mild ED	16.6%
Moderate	16.2%
Severe	9.4%
Mortality/hospice	5.9%

Table S1. NSAID use by COVID-19 severity among the unmatched COVID-19 positive cohort. NSAID use is likely incompletely captured disproportionately for outpatients and patients with less severe COVID-19, which may cause residual confounding.

Medication	Concept id
ibuprofen	1177480
ketorolac	1136980
diclofenac	1124300
celecoxib	1118084
indomethacin	1178663
naproxen	1115008
etodolac	1195492
lornoxicam	19049709
tenoxicam	19041220
piroxicam	1146810
droxicam	19056645
meloxicam	1150345
aspirin	1112807
acetaminophen	1125315

Table S2. OMOP concept IDs for medications. For each medication, the concept ID shown and all its descendants, excluding concepts representing topical and ophthalmic preparations, were used for analysis.

Condition	Concept id
Alcoholic liver damage	201612
Chronic hepatitis	200763 (excluded 4245975 and 201612)
Diabetes type 2	201826
Hepatic failure	4245975 (excluded 201612)
Hypertension	316866
Ischemic heart disease	4185932
Lupus	255891
Malignant neoplasm (lymphoid hematopoietic related tissue)	4147164
Neoplasm	438112 (excluded 4147164)
Nonischemic heart disease	321588 (excluded 4185932)
Vascular dementia	443605
Alzheimer's disease	378419 (excluded 443605)
Cerebral infarction	443454
Chronic respiratory disease	4063381
Dementia associated with another disease	374888 (excluded 378419 and 443605)
Diabetes type 1	201254
Hepatic fibrosis	4267417 (excluded 201612, 4245975, and 200763)
Hepatic steatosis	4059290 (excluded 201612, 4245975, 200763, and 4267417)
Hypertensive kidney disease	44782429
Nicotine dependence	4209423
Nonhypertensive chronic kidney disease	46271022 (excluded 44782429)
Other liver disease	194984 (excluded 201612, 4245975, 200763, 4267417, 4059290, 200451, and 192680)
Portal hypertension	192680
Rheumatoid arthritis	80809 (excluded 36684997)
Unspecified dementia	4182210 (excluded 443605, 378419, and 374888)
Psoriasis	140168

Table S3. OMOP concept IDs for conditions. For each condition, the concept ID shown and all its descendants were used for analysis.

Study	Cohort (n)	On NSAIDs	Main findings
Abu Esba 2021 [1]	503	146	No association between ibuprofen or any other NSAID and worse COVID-19 outcomes
Alamdari 2020 [2]	459	37	NSAID use had no substantial impact on mortality.
Bruce 2020 [3]	1222	54	No association between prior NSAID use and time to mortality or length of stay.
Chandan2021 [4]	25,659	13,202	No increase in mortality among patients with osteoarthritis in a primary care setting.
Choi 2020 [5]	293	8	Ibuprofen was not a risk factor associated with disease progression.
Drake 2021 [6]	78,674	4211	NSAID use was not associated with worse in-hospital mortality, critical care admission, requirement for invasive ventilation, requirement for non-invasive ventilation, requirement for oxygen, or occurrence of acute kidney injury.
Gianfrancesco 2020 [7]	531	111	NSAID use not associated with hospitalisation status.
Gupta 2020 [8]	2215	99	No significant association with 28-day mortality
Hwang 2020 [9]	103	5	NSAID use showed no statistically significant difference in death rate.
Imam 2020 [10]	1305	466	Patients using NSAIDs prior to hospitalization had lower odds of mortality.
Kragholm 2020 [11]	4002	264	No significant association between recent ibuprofen prescription claims and severe trajectory of COVID-19.
Lund 2020 [12]	9236	248	NSAIDs not associated with 30-day mortality, hospitalization, ICU admission, mechanical ventilation, or renal replacement therapy.
Park 2021 [13]	7590	398	NSAID use not associated with mortality or ventilator care in Covid-19 patients.
Rinott 2020 [14]	403	87	Ibuprofen use not associated with worse clinical outcomes, compared with paracetamol or no antipyretic.
Sahai 2021 [15]	911	465	NSAIDs use showed no significant differences in clinical covariates
Wong 2021 (#1) [16]	2,463,707	536,423	No evidence of difference in risk of COVID-19 related death associated with current use of NSAIDs
Wong 2021 (#2) [16]	1,708,781	175,495	In cohort of people with rheumatoid arthritis/osteoarthritis, a lower risk of COVID-19 related death was associated with current use of NSAID versus non-use.

Table S4. Previously published studies on potential associations of NSAID use with outcome in COVID-19.

S1 Supplemental Note 1

This project was conducted in the National Institute of Health (NIH) N3C Data Enclave on the Palantir Foundry platform (Palantir Technologies Inc., Denver, Colorado). This platform organizes code in nodes (transformations) that form a directed acyclic graph and cannot easily be presented in linearized fashion. The purpose of this documentation is to illustrate the approach taken by our analysis and to promote reproducibility and extensibility on the Palantir platform [17, 18] or (with appropriate changes) on other platforms.

The Palantir platform enables each node to use a different programming language. We chose SQL, Python, and R according to which technique was best adapted to the task at hand. Spark SQL (version 3.0.2-palantir.24), Python version 3.6.10, and R version 3.5.1 (2018-07-02) were used.

The N3C Data Enclave (hereafter “Enclave”) contains data from over 2.4 million COVID-19 positive patients from 64 health systems in the United States. The dataset utilized in this study was frozen on October 5, 2021. Data were harmonized, integrated, and mapped with the Observational Medical Outcomes Partnership (OMOP) 5.3.1 vocabulary.

For clarity, we have omitted most of the code that was used to check and output results and concentrate on the code that performed the analysis.

S1.1 Inputs

The inputs for our analysis included OMOP tables as well as some tables provided by the Palantir platform with processed data.

S1.1.1 OMOP Tables

The following OMOP tables were used for our analysis (Table S5). We refer the reader to the original OMOP documentation for more information details [19].

Table	Summary
drug_era	Represents a span of time when a patient is assumed to be exposed to a particular drug (active ingredient).
condition_occurrence	Dates when a condition is considered to have started and (if applicable) ended

Table S5. OMOP tables used in this study.

S1.1.2 Tables offered by the Enclave

The following Enclave-specific tables were used for our analysis.

S1.1.3 concept set members

This table defines the relations between the codesets and concept sets (Table S6).

codeset_id	concept_id	concept_set_name	concept_name
837398380	436540	Fracture	Open fracture axis
837398380	436824	Fracture	Open fracture of cervical vertebra without spinal cord injury
837398380	436826	Fracture	Closed fracture of shaft of radius
837398380	436832	Fracture	Closed fracture of sternum
837398380	437117	Fracture	Open fracture of intracapsular section of femur
(...)	(...)	(...)	(...)

Table S6. concept_set_members table. Some examples from this table, which is used to coordinate codesets created by N3C Enclave members (often using the OMOP/OHDSI Atlas tool). The table has three additional columns (not shown here) that specify the version and were used to extract the latest versions for analysis.

S1.2 complete patient table with derived scores

This table is prepared as described [18], and contains derived information that was used to provide information about some of the covariates used in our analysis (Table S7).

Column	data type	comment
person_id	string	unique patient identifier
data_partner_id	integer	identifier of N3C contributing center
visit_concept_id	integer	e.g., 9201 (Inpatient Visit)
visit_start_date	Date	The date of the initial medical encounter for COVID-19 or control condition
visit_concept_name	string	concept name corresponding to visit_concept_id
visit_occurrence_id	string	identifier for a visit
AKI_in_hospital	string	“Yes” or null
ECMO	string	“ECMO” or null
Invasive_Ventilation	string	“Invasive Ventilation” or null
positive_covid_test	string	“true” or null
negative_covid_test	string	“true” or null
Suspected_COVID	string	“true” or null
in_death_table	string	“true” or null. True if patient known to be deceased
age_at_visit_start_in_years_int	integer	age in years
length_of_stay	integer	length of stay in days
Race	string	e.g., “Missing/Unknown”
Ethnicity	string	e.g., “Missing/Unknown”
gender_concept_name	string	e.g., “FEMALE”
smoking_status	string	e.g., “Non smoker”
blood_type	string	e.g., “Unknown”
covid_status_name	string	e.g., “covid_confirmed_positive”
Severity_Type	string	e.g., “Mild”
InpatientOrED	boolean	true or false
Q_Score	integer	Charlson Comorbidity Index
BMI	double	e.g., 18.2
Height	double	in meters
Weight	double	in kilograms
Testcount	integer	Number of COVID tests performed

Table S7. Fields of the complete_patient_table_with_derived_scores table

S1.2.1 complete_patient_table_with_covariates

Inputs:

- complete_patient_table_with_derived_scores (Section S1.2)
- covariate_list. This is a table with true or false values (in columns) for the covariates that is derived from the condition_occurrence table and the corresponding concept sets in a straightforward fashion that is omitted here for conciseness.

```
SELECT *  
FROM complete_patient_table_with_derived_scores  
LEFT JOIN covariate_list USING(person_id)
```

Listing 1: Join the covariates to the patient table.

S1.3 create cohort dataset

Inputs:

- concept_set_members (Sec. S1.1.3)
- drug_era (Sec. S1.1.1)
- complete_patient_table_with_covariates (Sec. S1.2.1)

The following code creates the cohort dataset we will use for statistical analysis.

```
def create_cohort_dataset( concept_set_members , drug_era ,  
complete_patient_table_with_covariates ) -> None:  
    dat = complete_patient_table_with_covariates  
    covariates_to_check = [ "alcoholic_liver_damage" , "chronic_hepatitis" , "diabetes2" ,  
"hepatic_failure" , "hypertension" , "ischemic_heart_disease" , "lupus" ,  
"malignant_neoplasm_lymphoid_hematopoietic_related_tissue" ,  
"neoplasm" , "nonischemic_heart_disease" , "vascular_dementia" , "alzheimers_disease" ,  
"cerebral_infarction" , "chronic_resp" , "dementia_associated_with_another_disease" ,  
"diabetes1" , "hepatic_fibrosis" , "hepatic_steatosis" , "hypertensive_kidney_disease" ,  
"nicotine_dependence" , "nonhypertensive_chronic_kidney_disease" , "other_liver_disease" ,  
"portal_hypertension" , "rheumatoid_arthritis" , "unspecified_dementia" , "psoriasis" ,  
"immunosuppression" ]  
  
    min_num_of_cases_to_keep_covariate = 1000 # any cov with less than 1000 True's will be  
dropped  
  
    drug_concept_sets = [ "ibuprofen-107" , "oxicams-jac" , "ketorolac-9" , "diclofenac-28" , "  
celecoxib-2" , "indomethacin-72" , "naproxen-4" , "etodolac-36" ]  
    exclude_drug_concept_sets = [ "aspirin-1-9" , "acetaminophen-1-26" ]  
  
    # filter for COVID+  
    patient_table = dat.filter( dat.positive_covid_test==True )  
    drug_concept_ids = None  
  
    for this_concept_set_name in drug_concept_sets:  
        try:  
            these_drug_concept_ids = get_codeset( this_concept_set_name ,  
concept_set_members , get_is_most_recent=True , exclude_topicals=True ,  
verbose=verbose )  
        except NoConceptIdsError:  
            raise Exception( "I couldn't find any concept IDs for {this_concept_set_name}" )  
  
        if not drug_concept_ids:  
            drug_concept_ids = these_drug_concept_ids
```

```

34     else:
35         drug_concept_ids = DataFrame.unionAll(drug_concept_ids, these_drug_concept_ids
36 )
37
38 # make column with on-drug
39 pts_on_drug = get_pts_on_drug(patient_table, drug_era,
40 drug_concept_ids, verbose=False)
41 pts_on_drug = pts_on_drug.withColumnRenamed("person_id", "drug_person_id")
42 patient_table = patient_table.join(pts_on_drug,
43 patient_table.person_id == pts_on_drug.drug_person_id, how="leftouter")
44 patient_table = patient_table.withColumn("on_drug",
45 (patient_table.drug_person_id.isNotNull()))
46 patient_table = patient_table.drop("drug_person_id")
47 treated_pt_count = patient_table.filter(patient_table.on_drug==True).count()
48
49 # Flag patients taking other NSAIDs (e.g. acetaminophen, aspirin)
50 excluded_drug_concept_ids = None
51 for this_concept_set_name in exclude_drug_concept_sets:
52     # get drug concept IDs
53     try:
54         these_excluded_drug_concept_ids = get_codeset(this_concept_set_name,
55 concept_set_members, get_is_most_recent=True, exclude_topicals=True,
56 verbose=verbose)
57     except NoConceptIdsError:
58         raise Exception("(Excluded drug) I couldn't find any concept IDs for {
59 this_concept_set_name}")
60     if not excluded_drug_concept_ids:
61         excluded_drug_concept_ids = these_excluded_drug_concept_ids
62     else:
63         excluded_drug_concept_ids = DataFrame.unionAll(excluded_drug_concept_ids,
64 these_excluded_drug_concept_ids)
65
66 pts_on_excluded_drug = get_pts_on_drug(patient_table, drug_era,
67 excluded_drug_concept_ids, verbose=False)
68
69 # get list of pts on excluded drug and not on_drug of interest
70 not_on_drug_pts = patient_table.filter(patient_table.on_drug ==
71 False).select("person_id")
72 not_on_drug_pts = not_on_drug_pts.withColumnRenamed("person_id", "drug_person_id")
73
74 exclude_patients = pts_on_excluded_drug.join(not_on_drug_pts, pts_on_excluded_drug.
75 person_id == not_on_drug_pts.drug_person_id, how="inner").select("drug_person_id")
76
77 # make a new column with info about on_excluded_drug, so we can analyze cohort numbers
78 coherently in the next node
79 patient_table = patient_table.join(exclude_patients, patient_table.person_id ==
80 exclude_patients.drug_person_id, how="leftouter")
81 patient_table = patient_table.withColumn("on_excluded_drug", (patient_table.
82 drug_person_id.isNotNull()))
83 patient_table = patient_table.drop("drug_person_id")
84
85 # Some data cleaning on covariates
86 # merge rheumatoid-arthritis-w-factor and rheumatoid-arthritis columns
87 from pyspark.sql.functions import when
88 patient_table = patient_table.withColumn("ra_or_ra_with_factor", when(patient_table.
89 rheumatoid_arthritis_w_factor == True, True)
90 .when(patient_table.rheumatoid_arthritis == True, True)
91 .otherwise(False))
92 patient_table = patient_table.drop("rheumatoid_arthritis", "

```

```

86  rheumatoid_arthritis_w_factor")
    patient_table = patient_table.withColumnRenamed("ra_or_ra_with_factor", "
rheumatoid_arthritis")

88  # replace null's with False
    patient_table = patient_table.fillna(value=False, subset=covariates_to_check)

90  # remove covariates with less than 1000 patients
92  for cov in covariates_to_check:
    ncases = patient_table.filter(patient_table[cov] == True).count()
94    if ncases < min_num_of_cases_to_keep_covariate:
        print(f"dropping {cov} column since it's less than
min_num_of_cases_to_keep_covariate ({min_num_of_cases_to_keep_covariate})")
96        patient_table = patient_table.drop(cov)

98  return(patient_table)

```

Listing 2: Create the case/control dataset

S1.4 drop outpatient and ed

Inputs:

- `create_cohort_dataset` (S1.3; note that we omit an intermediate step that removes data from centers that do not provide BMI data)

This node performs drops outpatients as well as outpatients who were seen in an emergency department (ED) but not admitted.

In effect, this node drops visits classified as “Outpatient Visit”, “Emergency Room Visit”, “No matching concept”, “Office Visit”, “Information not available”, “Telehealth”, “Non-hospital institution Visit”, “Observation Room”, “Ambulatory Surgical Center”, “Ambulatory Clinic / Center”, “Laboratory Visit”, “Interactive Telemedicine Service”, “Ambulatory Radiology Clinic / Center”, “Ambulatory Infusion Therapy Clinic / Center”, “Ambulatory Dental Clinic / Center”, “Ambulatory Oncology Clinic / Center”, “Ambulatory Magnetic Resonance Imaging (MRI) Clinic / Center”, “Ambulatory Mammography Clinic / Center”, “Ambulatory Oncological Radiation Clinic / Center”, “Home Visit”, “Ambulatory Endoscopy Clinic / Center”, “Skilled Nursing Facility”.

```

drop_outpatient_and_ed <- function(create_cohort_dataset) {
2  dat <- throw_out_data_partners_with_100_perc_missing_BMI
    dat <- dat[dat$visit_concept_name %in% c("Inpatient Visit", "Inpatient Hospital",
4  "Emergency Room and Inpatient Visit", "Inpatient Critical Care Facility" ),]
    return(dat)
6  }

```

Listing 3: Remove outpatients

S1.5 do matchit drop outpatient and ed

Inputs:

- `drop_outpatient_and_ed` (Section S1.4)

This node performs propensity matching on data that contains only inpatients (outpatient and outpatient emergency department have been dropped).

Todo – explain `on_excluded_drug`

```

do_matchit_drop_outpatient_and_ed <- function(drop_outpatient_and_ed) {
2   covariates_to_exact_match <- NULL
   excluded_drug_col <- "on_excluded_drug"
4   dat <- drop_outpatient_and_ed

6   # cols we need below
   factor_cols <- c("Race", "Ethnicity", "gender_concept_name", "smoking_status",
8     "InpatientOrED", "Severity_Type", "data_partner_id")
   outcomes <- c("Severity_Type", "in_death_table", "Invasive_Ventilation",
10    "AKI_in_hospital", "ECMO", "InpatientOrED")
   treatment <- "on_drug"
12  data_partner_col = "data_partner_id"
   patient_info_for_matching <- c("age_at_visit_start_in_years_int", "Race", "Ethnicity",
14    "gender_concept_name", "smoking_status", "BMI", "Q_Score")
   comorbidities_for_matching <- c("alcoholic_liver_damage", "chronic_hepatitis",
16    "diabetes2", "hepatic_failure", "hypertension", "ischemic_heart_disease",
18    "lupus", "malignant_neoplasm_lymphoid_hematopoietic_related_tissue",
20    "neoplasm", "nonischemic_heart_disease", "vascular_dementia",
22    "alzheimers_disease", "cerebral_infarction", "chronic_resp",
24    "dementia_associated_with_another_disease", "diabetes1",
    "hepatic_fibrosis", "hepatic_steatosis", "hypertensive_kidney_disease",
    "nicotine_dependence", "nonhypertensive_chronic_kidney_disease",
    "other_liver_disease", "portal_hypertension", "rheumatoid_arthritis",
    "unspecified_dementia", "psoriasis")

26  # filter out patients "on_excluded_drug"
   dat <- filter(dat, dat[[excluded_drug_col]] == FALSE)

28

   uniq_pts = length(unique(dat$person_id))
30  print(paste("found", uniq_pts, "person_ids after filtering out pts on_excluded_drug"))

32  # clean/convert outcome columns into boolean
   dat$in_death_table = !is.na(dat$in_death_table) & dat$in_death_table == TRUE
34  dat$Invasive_Ventilation = !is.na(dat$Invasive_Ventilation)
   dat$ECMO = !is.na(dat$ECMO)
36  dat$AKI_in_hospital = !is.na(dat$AKI_in_hospital)

38  cols_of_interest <- c(
   # fairly standard patient info
40    "person_id", # always keep this
   outcomes,
42    treatment,
   data_partner_col,
44    patient_info_for_matching,
   comorbidities_for_matching)

46

   matchit_formula <- as.formula(paste(treatment, "~",
48     paste(c(patient_info_for_matching, comorbidities_for_matching, data_partner_col),
        collapse="+")))
   matched_data <- do_matchit(data=dat, cols_of_interest=cols_of_interest,
50     matchit_formula=matchit_formula, factor_cols=factor_cols,
52     matchit_method="nearest", verbose=verbose,
   exact_matching=covariates_to_exact_match, do.love=TRUE, make_table_one=FALSE,
54     vim.margin.plot.cols=c("age_at_visit_start_in_years_int", "BMI"))

56  if (!exists("matched.data") || is.null(matched.data)){
   stop("matched.data is null after matching")
58  }

60  return(matched.data)

```

```
}
```

Listing 4: propensity matching

S1.6 glm regression inpatient no ed mild mild ed moderate versus severe dead

Inputs:

- `do_matchit_drop_outpatient_and_ed` (Section S1.5)

The following code expects as input a data table with matched patients and controls. It uses a convenience function called `set_cols_as_factors` that converts columns to factor using the following command for each of the selected columns.

```
dat[[x]] <- as.factor(dat[[x]])
```

```
glm_regression_inpatient_no_ed_mild_mild_ed_moderate_versus_severe_dead <- function(do_
  matchit_drop_outpatient_and_ed) {
2   matched.data <- do_matchit_drop_outpatient_and_ed
   factor_cols <- c("Race", "Ethnicity", "gender_concept_name", "smoking_status",
4   "InpatientOrED", "Severity_Type", "data_partner_id")
   glm_outcomes <- c("Severity_Type", "in_death_table", "Invasive_Ventilation",
6   "AKI_in_hospital", "ECMO")

8   matched.data <- set_cols_as_factors(matched.data, factor_cols, verbose=FALSE)

10  matched.data[["Severity_Type"]] <- as.numeric(matched.data[["Severity_Type"]] %in% c("
  Severe", "Dead_w_COVID"))

12  # these were selected by the code in "select_covariates" node
   covariates_always_keep <- c("on_drug", "age_at_visit_start_in_years_int", "Race",
14  "Ethnicity", "gender_concept_name", "smoking_status", "BMI")
   selected_covariates <- c("alcoholic_liver_damage",
16   "alzheimers_disease",
   "cerebral_infarction",
18   "chronic_resp",
   "diabetes1",
20   "diabetes2",
   "hepatic_failure",
22   "hepatic_fibrosis",
   "hypertensive_kidney_disease",
24   "ischemic_heart_disease",
   "lupus",
26   "malignant_neoplasm_lymphoid_hematopoietic_related_tissue",
   "neoplasm",
28   "nicotine_dependence",
   "nonhypertensive_chronic_kidney_disease",
30   "nonischemic_heart_disease",
   "other_liver_disease",
32   "portal_hypertension",
   "psoriasis",
34   "Q_Score",
   "unspecified_dementia",
36   "vascular_dementia")

38  covariate_terms_for_formula <- paste(c(covariates_always_keep, selected_covariates),
  collapse="+")
```

```

40 # do logistic regression
42 all_glms <- lapply(glm_outcomes, function(outcome){
44     logit.formula <- as.formula(paste(outcome, "~", covariate_terms_for_formula))
46     logit <- NULL
48     try(logit <- do_glm(dat=matched.data, glm.formula=logit.formula))
50     return(logit)
52 })
54 print(all_glms)
56
58 # make table with summary
60 parsed_data <- lapply(all_glms, function(logit){
62     logit_summary <- summary(logit)
64     logit_coef <- coef(logit_summary)
66     all_coeff_data <- cbind(logit_coef, data.frame("OR"=exp(logit_coef[, 'Estimate']),
68         exp(confint(logit)))
70     all_coeff_data <- all_coeff_data["on_drugTRUE", ] %>%
72     dplyr::select(-c("Estimate", "Std. Error", "z value"))
74     return(all_coeff_data)
76 })
78 print(cbind(glm_outcomes, do.call(rbind, parsed_data)))
80
82 return(NULL)
84 }

```

Listing 5: glm regression.

S1.7 generate outcome counts table inpatients no ed

Inputs:

- do_matchit_drop_outpatient_and_ed (Section S1.5)

This script is used to generate output as such as the following.

```

# A tibble: 3 x 6
# Groups:   on_drug [1]
  on_drug Severity_Type      n perc   n1 perc1
  <lgl>   <ord>          <int> <dbl> <int> <dbl>
1 TRUE   Moderate      18023 91.3  16972 86.0
2 TRUE   Severe         776 3.93  1047 5.30
3 TRUE   Dead_w_COVID  947 4.80  1727 8.75
# A tibble: 4 x 5
  outcome          n perc   n1 perc1
  <chr>          <int> <dbl> <int> <dbl>
1 in_death_table  947 4.80  1727 8.75
2 Invasive_Ventilation 1150 5.82  1867 9.46
3 AKI_in_hospital  1729 8.76  2437 12.3
4 ECMO            54 0.273  109 0.552

```

```

generate_outcome_counts_table_inpatients_no_ed <-
2 function(do_matchit_drop_outpatient_and_ed) {
4     matched.data <- do_matchit_drop_outpatient_and_ed
6     secondary_outcomes <- c("Severity_Type", "in_death_table", "Invasive_Ventilation",
8         "AKI_in_hospital", "ECMO")
10    factor_cols <- c("Race", "Ethnicity", "gender_concept_name", "smoking_status",

```

```

8   "InpatientOrED", "Severity_Type", "data_partner_id")
matched.data <- set_cols_as_factors(matched.data, factor_cols, verbose=FALSE)
matched.data <- fix_ordinal_outcome(matched.data)

10
12  summaries <- lapply(secondary_outcomes, function(this.outcome){
    this.outcome.tab <- matched.data %>% dplyr::group_by(on_drug) %>% dplyr::count(!!
sym(this.outcome))
    this.outcome.tab <- this.outcome.tab %>% dplyr::mutate(perc = 100 * n / sum(n))
14  this.outcome.tab <- cbind(this.outcome.tab %>% filter(on_drug), this.outcome.tab
%>% filter(!on_drug))
    return(this.outcome.tab)
16  })

18  # severity type data:
std <- summaries[[1]] %>% dplyr::select(Severity_Type, n, perc, n1, perc1)
20  print(std)

22  secondary_outcomes <- lapply(summaries[2:5], function(dat){
    # getting rid of the "FALSE" row because it's unnecessary, and using the name of
the 2nd column as the outcome
24  dat <- dat[dat[,2]==TRUE,] %>% mutate(outcome=colnames(dat)[2]) %>% dplyr::select(
outcome, n, perc, n1, perc1) %>% ungroup() %>% dplyr::select(-on_drug)
    return(dat)
26  })
28  print(dplyr::bind_rows(secondary_outcomes))
return(NULL)
}

```

Listing 6: Generate outcome counts.

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