



Published in final edited form as:

Genet Med. 2021 April ; 23(4): 758–766. doi:10.1038/s41436-020-01038-0.

Health and economic outcomes of newborn screening for infantile-onset Pompe disease

John S. Richardson, PhD^{1,2}, Alex R. Kemper, MD, MPH³, Scott D. Grosse, PhD⁴, Wendy K. K. Lam, PhD⁵, Angela M. Rose, MPH⁶, Ayesha Ahmad, MD⁷, Achamyeleh Gebremariam, MS⁶, Lisa A. Prosser, PhD^{1,6,✉}

¹Department of Health Management and Policy, University of Michigan, Ann Arbor, MI, USA.

²RTI International, Research Triangle Park, NC, USA.

³Division of Ambulatory Pediatrics, Nationwide Children's Hospital, Columbus, OH, USA.

⁴Centers for Disease Control and Prevention, Atlanta, GA, USA.

⁵Duke Clinical & Translational Science Institute, School of Medicine, Duke University, Durham, NC, USA.

⁶Susan B. Meister Child Health Evaluation and Research (CHEAR) Center, Department of Pediatrics, University of Michigan, Ann Arbor, MI, USA.

⁷Division of Pediatric Genetics, Department of Pediatrics, University of Michigan, Ann Arbor, MI, USA.

Abstract

PURPOSE: To estimate health and economic outcomes associated with newborn screening (NBS) for infantile-onset Pompe disease in the United States.

METHODS: A decision analytic microsimulation model simulated health and economic outcomes of a birth cohort of 4 million children in the United States. Universal NBS and treatment was compared with clinical identification and treatment of infantile-onset Pompe disease. Main outcomes were projected cases identified, costs, quality-adjusted life-years (QALYs), and incremental cost-effectiveness ratios (ICERs) over the life course.

Reprints and permission information is available at <http://www.nature.com/reprints>

[✉]**Correspondence** and requests for materials should be addressed to L.A.P. lisapros@umich.edu.

AUTHOR CONTRIBUTIONS

Conceptualization: J.S.R., A.R.K., S.D.G., and L.A.P. Data curation: A.G. Formal analysis: J.S.R., A.G., and L.A.P. Funding acquisition: L.A.P. Investigation: J.S.R., A.R.K., S.D.G., W.K.K.L., A.M.R., A.A., A.G., and L.A.P. Methodology: J.S.R., A.G., and L.A.P. Project administration: A.M.R. and L.A.P. Supervision: L.A.P. Visualization: A.M.R. and A.G. Writing-original draft: J.S.R., S.D.G., and L.A.P. Writing-review & editing: J.S.R., A.R.K., S.D.G., W.K.K.L., A.M.R., A.A., A.G., and L.A.P.

DATA AVAILABILITY

The model is available upon request by contacting the corresponding author.

CODE AVAILABILITY

The model is available upon request by contacting the corresponding author.

COMPETING INTERESTS

The authors declare no competing interests.

The online version of this article (<https://doi.org/10.1038/s41436-020-01038-0>) contains supplementary material, which is available to authorized users.

RESULTS: Universal NBS for Pompe disease and confirmatory testing was estimated to cost an additional \$26 million annually. Additional medication costs associated with earlier treatment initiation were \$181 million; however, \$8 million in medical care costs for other services were averted due to delayed disease progression. Infants with screened and treated infantile-onset Pompe disease experienced an average lifetime increase of 11.66 QALYs compared with clinical detection. The ICER was \$379,000/QALY from a societal perspective and \$408,000/QALY from the health-care perspective. Results were sensitive to the cost of enzyme replacement therapy.

CONCLUSION: Newborn screening for Pompe disease results in substantial health gains for individuals with infantile-onset Pompe disease, but with additional costs.

INTRODUCTION

State-based newborn screening (NBS) programs have expanded over the past 50 years from screening for phenylketonuria to screening for more than 30 conditions.^{1,2} The Advisory Committee on Heritable Disorders in Newborns and Children (ACHDNC) was developed to support state NBS decisions by reviewing proposed conditions and synthesizing evidence regarding the feasibility and benefits of screening.³ The ACHDNC makes recommendations to the Secretary of Health and Human Services, who in turn determines a Recommended Uniform Screening Panel (RUSP); cost-effectiveness is not a criterion in the recommendation process.⁴

Pompe disease was recommended by the ACHDNC in June 2013 and was added to the RUSP in March 2015.⁵ It is a rare condition that is identified in about 1 in 40,000 births.⁶ Pompe disease occurs from a defect in the *GAA* gene leading to the accumulation of lysosomal glycogen and, depending on the form and severity, can result in cardiomyopathy, progressive muscle weakness, respiratory failure, and heart failure.⁷ Pompe disease can be treated with alglucosidase alfa, an enzyme replacement therapy (ERT) that costs between \$250,000 and \$500,000 per year for biweekly infusions.⁸

The purpose of this study was to evaluate the cost-effectiveness of universal NBS for infantile-onset Pompe disease compared with usual clinical identification.

MATERIALS AND METHODS

Study design and scope

We created a decision analytic microsimulation model for a US cohort of 4 million newborns.⁹ Microsimulations are mathematical models that track the unique health trajectory and health outcomes of each simulated person over time. The model compared outcomes for two strategies: (1) clinical identification of infantile-onset Pompe disease and treatment with ERT; and (2) NBS for Pompe disease and treatment of infantile-onset disease with ERT. The analytic time horizon included the life course of the birth cohort. We used one-year time increments to capture changes in health status over each individual's life. We incorporated costs and outcomes from the health-care sector and societal perspectives.¹⁰ Insufficient data are available to model the long-term costs and outcomes for individuals identified through NBS with late-onset Pompe disease.

Model structure and assumptions

In the clinical identification scenario, individuals were diagnosed with Pompe disease at varying ages based on testing prompted by the recognition of symptoms. This analysis used a classification of Pompe disease developed for the ACHDNC evidence review.^{4,11} The cases were classified as infantile-onset (those with onset before age 12 months) or late-onset. Infantile-onset cases were further classified as either with or without cardiomyopathy (Fig. 1). A more recent set of guidelines defines “infantile-onset” as only those with cardiomyopathy and onset before 12 months; however, these were published after our analytic framework had been defined.¹² Once identified as having Pompe disease, an individual in our analysis can remain at the current stage of disease severity, progress to later stages of symptoms, or die.

In the NBS scenario, we assume that newborns are screened within 2 days of birth. If a newborn has a positive screen, a repeat test is conducted, followed by, if indicated, confirmatory testing of enzyme activity and molecular testing and diagnosis by 2–3 weeks of age. Infants with low enzyme activity and *GAA* variants consistent with pseudodeficiency alleles are not evaluated further. Individuals diagnosed with Pompe disease are classified as infantile-onset (with or without cardiomyopathy) or probable late-onset depending on whether signs or symptoms are present during confirmatory testing. Those in the NBS scenario with infantile-onset Pompe progress through the same set of long-term health states as those in the clinical identification scenario. Probability of a false negative result is also included in the NBS scenario. It is assumed that all such cases would be late-onset.

The focus of our analysis is on infantile-onset cases and we did not evaluate changes in health outcomes or costs of treatment for late-onset Pompe disease. However, we did include an extra annual physician visit for asymptomatic patients screened and diagnosed with probable Pompe disease who had not yet exhibited symptoms as standard of care.

In the clinical identification scenario we assumed that identification of infantile-onset Pompe disease among those with cardiomyopathy occurred at 4.5 months of life.⁶ In the absence of screening, we assumed that one-third of those with infantile-onset Pompe disease but no cardiomyopathy were identified and initiated treatment earlier than one year, while two-thirds were identified after the first year of life with an average age of diagnosis at three years before being identified and starting treatment.

The life course trajectory of infantile-onset Pompe disease was captured using a state-transition simulation model with the following states: no symptoms, mild symptoms, severe symptoms, died from Pompe disease, and died from other causes (Fig. 1). This state-transition model was a part of the microsimulation model and tracks individuals as they cycle through the different health states over consecutive 1-year time periods. Mild symptoms included some signs and symptoms. Ventilator dependence is required for the classification of severe disease.

Model inputs were derived using published reports, unpublished data, and judgments from an expert panel (Table 1). The expert panel consisted of six individuals (see “Acknowledgements”) with extensive experience in newborn screening, studying rare

genetic conditions, and treating individuals with Pompe disease. The parameters determined from the expert panel were developed using a modified Delphi or “Decision Delphi” approach.¹³

Epidemiology inputs

The sensitivity and specificity of NBS as well as other screening probabilities were derived from a pilot screening program in Taiwan and confirmed by an expert panel (Table S2).¹⁴ Transition probabilities were calibrated to short-term health outcomes (to age 3) using published and unpublished data and longer-term outcome trajectories determined by an expert panel^{15,16} (Figure S2, Table S1). Based on input from the expert panel, we also assumed that early treated individuals with infantile-onset Pompe disease have the same underlying risk of neurological or other conditions as the general pediatric population.

Costs

Costs included the costs of screening all newborns, doing confirmatory testing among positive screens, and the costs associated with treatment for patients diagnosed with infantile-onset Pompe disease. Both medical costs and time costs for informal caregivers (for the societal perspective analysis) were included (Table S3). All costs were adjusted to 2016 dollars (Table S3). The cost of screening was determined through key informant interviews with US states that were either in the process of starting to implement or had established screening for Pompe disease. A base case estimate of \$6.40 per newborn was used for the cost of the initial screen (Table 1). Our cost estimates for false positive screens presume that programs use an assay with specificity comparable with that reported by the New York screening program rather than one of the less efficient assays.¹⁷ In the base case, we assumed full adherence to treatment recommendations and that no family would deny treatment for their child. Patients with confirmed infantile-onset Pompe disease were assumed to receive an infusion of 20 mg/kg every other week, resulting in medication costs that ranged from an estimated \$75,475 to \$513,233 per year depending on age and on the assumed cost of the drug in addition to administration costs of \$14,300 per year.

We used a microcosting approach to estimate the total costs of care associated with each health state (Table S3). A microcosting approach establishes estimates for each of the components of resource units and then aggregates them to get the overall costs. In addition to the costs of ERT, ongoing costs of care included costs of additional required medications, specialist visits, emergency visits, hospitalizations, laboratory tests, and medical equipment, as well as formal and informal caregiving (Table S3). The costs of these medical services were based on the Centers for Medicare and Medicaid Services Physician Fee Schedule, and the number of services were based on estimates from clinical experts (Table S3). Informal caregiving is care provided by a family member or other individual who is not paid for his or her time. Informal caregiving hours were assumed to range from 14 hours per week for patients with few symptoms to 16 hours per day for severe forms.¹⁸ Formal caregiving hours were assumed to be 8 hours per day for those with severe symptoms. Medical expenditures unrelated to Pompe disease were also included.¹⁹

Quality of life

Health utility weights for each health state were determined from a separate study using time tradeoff questions fielded to a national community-based sample.²⁰ The values for those with mild symptoms ranged from 0.799 to 0.853, and for those with severe symptoms ranged from 0.399 to 0.536, depending on age. In scenario analyses, we also included reductions in health utility for family members attributable to the patient's illness (family spillover utility) as well as a small and transient reduction in health utility associated with the experience of a false positive screen.

Analysis plan

The primary outcome is the incremental cost-effectiveness ratio (ICER) in dollars per quality-adjusted life-year (QALY) gained for the NBS scenario relative to the clinical identification scenario. Incremental costs include the costs of screening, short-term follow-up, and diagnostic testing for all infants who screen positive. Secondary outcomes included numbers of cases identified with and without screening, disaggregated outcomes (i.e., the major components contributing to the numerator and denominator of the ICER), and undiscounted outcomes (i.e., the outcomes in 30 years are assumed to have the same value as outcomes over the next year). The base case analysis used most likely point estimates for parameters, and included test characteristics and screening costs derived from the New York screening program. We conducted one-way sensitivity analyses of screening parameters, transition probabilities, and cost parameters to determine the effect of varying individual parameters within a plausible range on the incremental cost-effectiveness ratio. Scenario analyses were also conducted for a broader range of hypothetical reduction in the cost of ERT, the inclusion of family spillover effects on quality of life, and a comparison between clinical identification with and without ERT. Given that clinicians sometimes increase the dosage or frequency of ERT, we also examined scenario analyses in which patients received an ERT dose of 40 mg per kg every other week and every week (instead of 20 mg/kg every other week in base case).²¹ Costs and QALYs were discounted using a 3% annual discount rate. Simulation software was TreeAge Pro 2017 version R2.1 (TreeAge Software, Inc., Williamstown, MA).

RESULTS

Screening test outcomes and diagnoses

Using base case parameter input values, 100 individuals in the simulated birth cohort would be identified with Pompe disease in both scenarios. Among these individuals, 34 have infantile-onset Pompe disease with cardiomyopathy, 6 infantile-onset (atypical) Pompe disease without cardiomyopathy, and 60 have late-onset Pompe disease. Newborn screening is also assumed to lead to the identification of 38 individuals who have low α -glucosidase enzyme activity due to being heterozygous for a pathogenic variant or a pseudodeficiency in a *GAA* allele that does not cause clinical symptoms.

Health outcomes

Newborn screening resulted in substantial health gains as measured by the proportion of patients who are ventilator-dependent, Pompe disease-related deaths, and QALYs for those with infantile-onset Pompe disease (Table 2). The QALYs gained from NBS compared with clinical identification was 12.9 QALYs per person with infantile-onset Pompe disease with cardiomyopathy. For infantile-onset Pompe disease without cardiomyopathy (atypical presentation), the gain was 4.7 QALYs per person. For a cohort of 4 million children, NBS results in a cumulative lifetime gain of 466 QALYs compared with clinical identification.

Costs

The total discounted lifetime costs of care for infantile-onset Pompe disease were projected at \$7.98 million for each person who is clinically identified (Table 2). For infantile-onset Pompe disease identified through screening, average lifetime costs were estimated to be \$11.7 million. Increases in costs of care associated with the screened group were predominantly from additional time receiving ERT due to earlier initiation of treatment combined with longer overall period of treatment due to averted deaths. Increased treatment costs were only slightly offset by improved health, resulting in lower infantile-onset Pompe disease-related costs other than ERT (−\$0.2 million), formal caregiving costs (−\$0.26 million), and informal caregiving costs (−\$0.35 million).

At the population level, NBS and confirmatory tests for 4 million births were estimated to cost \$26 million (Table 2). Total incremental costs associated with NBS strategy for infantile-onset Pompe disease were \$177 million from the societal perspective and \$190 million from the health sector perspective. ERT costs were estimated to increase by \$181 million under NBS. The category of cost with the largest decrease attributable to NBS was informal caregiving costs (−\$14 million).

Cost-effectiveness

Using base case assumptions, the ICER for NBS compared with clinical identification of infantile-onset Pompe disease was \$379,000/QALY from a societal perspective and \$408,000/QALY from a health sector perspective (Table 2). If the cost of screening were \$17.50 per test rather than \$6.40 per test, NBS would cost society 25% more or \$474,000/QALY. Lower estimates for the cost of ERT resulted in a cost-effectiveness ratio of \$303,000/QALY and higher estimates resulted in \$402,000/QALY (Fig. 2, Table S4). Varying treatment effectiveness across the plausible range yields cost-effectiveness ratios from \$311,000/QALY to \$420,000.

Figure S1 and Table S5 show a scenario analysis of reductions in the cost per dose of alglucosidase alfa. In scenario analysis, if the cost of the drug were reduced by 80% (discount to average wholesale price), the cost-effectiveness ratio would be approximately \$102,000/QALY (Figure S1 and Table S5). Results were robust to changes in other parameters. Including a family spillover utility for one parent yielded slightly more favorable cost-effectiveness results of \$372,000/QALY gained (Table S6). Increasing the dosage of ERT to 40 mg per kg every other week or every week substantially increases the ICER to \$750,000/QALY to \$1,500,000/QALY (Table S7).

In the scenario analysis where we evaluated the cost-effectiveness of ERT by comparing outcomes of clinical identification with ERT to clinical identification without ERT (in the absence of newborn screening), cost-effectiveness ratios were greater than \$500,000/QALY for clinical identification and treatment of infantile-onset Pompe disease (Table S8).

DISCUSSION

We found that NBS for Pompe disease in the United States would provide substantial gains in health and quality of life among individuals with infantile-onset Pompe disease. This comes, however, with increased costs due to earlier initiation of treatment and extended lifespan for individuals with infantile-onset Pompe disease. We measured the value gained from these increased costs through cost-effectiveness analysis.^{22,23} In situations where resources are limited, unofficial thresholds for ICERs are sometimes used in the United States to encourage investment in high value or low cost per QALY services. An ICER of \$379,000/QALY is far beyond the upper bound of the unofficial range of commonly used cost-effectiveness benchmarks (\$50,000–\$150,000/QALY) for the United States.²⁴ Some, however, argue that different criteria should be applied for the cost-effectiveness of treatments for rare conditions. The UK National Institute for Health and Clinical Excellence (NICE) for “highly specialized technologies” for conditions with a prevalence of 2 per 100,000 or less sets a cost-effectiveness threshold that is roughly three times higher than the general threshold.^{25,26} If a similar approach were applied in the United States to interventions for very rare disorders (i.e., a threshold range of \$150,000 to \$450,000/QALY), screening for infantile-onset Pompe disease would fall at the very upper end of what could be considered cost-effective using these alternate criteria. Other countries also cite additional considerations for treatments for rare diseases but have not put forth an adjusted threshold.²⁵ If higher dosages and frequency of ERT are delivered than the base case scenario, then newborn screening for infantile-onset Pompe disease would not meet thresholds for cost-effectiveness even using the most liberal criteria for assessing the cost-effectiveness of rare conditions.

This analysis extends previous work conducted on behalf of the Condition Review Workgroup for the Advisory Committee on Heritable Disorders in Newborns and Children.^{4,6} It evaluates the cost-effectiveness of NBS for infantile-onset Pompe disease, including costs and outcomes throughout the lifespan. In a prior analysis, we estimated that NBS in the United States would prevent 13 deaths and 26 people from needing mechanical ventilation by age 3 years. The current analysis extends these results into a lifetime simulation model to answer key questions on lifetime health and quality-of-life benefits while also incorporating screening and disease-related costs.

This analysis uses the strategy of clinical identification followed by treatment with alglucosidase alfa as the primary comparator strategy, recognizing that individuals identified clinically are being offered and typically will initiate treatment with alglucosidase alfa. In a scenario analysis comparing clinical identification with ERT to clinical identification without ERT, we estimated a much higher cost-effectiveness ratio. Even if higher thresholds are considered, this leaves us in a challenging position in which NBS for a condition might

potentially be considered cost-effective but the treatment for the condition at >\$500,000/QALY would not be cost-effective even under alternate criteria.

Two studies have assessed the cost-effectiveness of providing ERT, separate from NBS for Pompe disease.^{27,28} In the Netherlands, using a societal perspective, Kanters et al. estimated that alglucosidase alfa treatment of Pompe disease compared with no treatment was greater than €1 million/QALY.²⁷ A study from a health sector perspective estimated a cost of £234,308/QALY in England but did not include outpatient medical costs.²⁸ Neither study assessed the cost-effectiveness of NBS for Pompe disease. Our results for the cost-effectiveness of treatment with ERT for clinically identified infantile-onset Pompe disease appear broadly consistent with these studies.

Our results were not sensitive to the costs of screening. Determining the costs of NBS for a single condition can be difficult if screened for as part of a panel of lysosomal storage disorders. If costs of screening for Pompe disease as part of a panel were just \$1, the ICER would still be \$333,000/QALY, which is substantially greater than conventional cost-effectiveness thresholds. In the future, better information on costs of screening from NBS programs could improve the accuracy of cost-effectiveness analyses, although that may matter more for conditions with relatively low treatment costs.

In contrast, the results were sensitive to the cost of treatment. The high cost of treatments for rare conditions has been the subject of increasing debate.²⁹ Reducing the costs of ERT would result in more favorable cost-effectiveness ratios for NBS. If the cost of alglucosidase alfa were reduced by 80%, the cost-effectiveness ratio for NBS for infantile-onset Pompe disease would fall below \$150,000/QALY.

Inclusion of changes in quality of life for family members of a child with infantile-onset Pompe (family spillover effects) did not substantially affect the cost-effectiveness ratios. While the earlier initiation of ERT can improve the quality of life for children with Pompe disease, it does not eliminate spillover; and as lifespan increases, the duration of spillover also increases.

The use of a societal perspective instead of a narrower health-care sector perspective had a modest effect on the ICER remaining within the same order of magnitude as the ICER using a health-care sector perspective. The additional costs included in the societal analysis were informal caregiving and patient time costs, which, while substantial, were modest relative to the cost of ERT (Supplemental Table S11, Impact Inventory). Since Pompe disease is not associated with cognitive impairment, special education costs would not be applicable. Although productivity costs are substantial, they are offset by reduced consumption costs; guidance for the inclusion of both categories for pediatric populations is not currently available.

The evidence base for Pompe disease and other rare conditions is scarce. Lack of long-term outcomes data for treated patients required us to rely on expert judgment to estimate health trajectories for treated patients. Evidence on costs of living with Pompe disease was also sparse. Despite the uncertainty in these parameters, our results were robust except to changes in the costs of care for infantile-onset Pompe disease. Due to the complexity of the

disease model and the size of the modeled cohort, it was not feasible to conduct probabilistic sensitivity analysis in the study due to computational limitations.

The use of Medicare fee schedules is a commonly used approach by cost-effective analysts for costing health service utilization.¹⁰ In recent years, the Centers for Medicare and Medicaid Services have not increased physician fees for many services and this could result in a potential limitation for this study. If costs of medical services are underestimated, this would underestimate the benefits of screening and earlier treatment, yielding less favorable ratios than if costs of medical care were higher. However, since ERT represents the majority of the health-care costs for treated individuals, the impact of this limitation would be limited.

Newborn screening for infantile-onset Pompe disease also has implications for identification and treatment of late-onset Pompe disease, but these were not captured in this study to be consistent with the ACHDNC process to evaluate the benefits and harms of conditions that would be addressed during the newborn and childhood ages (typically the first year of life).³ The evidence review conducted for the ACHDNC focused on infantile-onset Pompe disease, consistent with the policy question faced by ACHDNC.⁶ Our analysis does include the costs of following probable late-onset cases but does not include any benefits or costs associated with treatment with ERT. One study has found ERT to improve the quality of life of those with late-onset Pompe during the first year of treatment, and to stabilize the quality of life during the subsequent years of treatment;³⁰ however, treatment protocols for the use of ERT in late-onset patients are still being established. Newer treatment guidelines suggest that early initiation of ERT is beneficial for some patients with late-onset Pompe disease, particularly those with less-advanced disease.³¹ Genetic testing following newborn screening can help identify individuals who are most likely to experience late-onset Pompe and eventually need ERT. Some pathogenic *GAA* variants are strongly correlated with late-onset and not infantile-onset Pompe disease.³² Since newborn screening is likely to result in regular clinical follow-up for individuals identified through screening but not diagnosed with infantile-onset Pompe disease, this will likely lead to earlier initiation of ERT for late-onset Pompe disease as a result of newborn screening. As more data become available, future studies should evaluate the cost-effectiveness of early diagnosis and treatment for late-onset Pompe disease. However, because of the cost of monitoring individuals with late-onset Pompe disease over many years and because the benefit of diagnosis may not occur for decades, including late-onset Pompe disease would lead to a less favorable cost-effectiveness assessment. Unfortunately, there are not sufficient data to model the inclusion of late-onset Pompe disease in this microsimulation study.

Newborns with pseudodeficiency may also experience additional costs following identification during NBS even though pseudodeficiency is not associated with any adverse health outcomes. Studies in Canada found modestly higher health-care utilization by children who screened positive for certain conditions but did not receive a diagnosis.^{33,34} Future studies evaluating long-term outcomes from NBS for Pompe disease could consider collecting data on this group of infants as well.

Conclusions

Newborn screening for infantile-onset Pompe disease yields considerable health gains and may be cost-effective in the context of rare diseases. As other rare heritable disorders are considered for NBS, this study provides insights into the relative costs and health benefits of screening. First, NBS may be considered cost-effective relative to no screening even when the underlying treatment is not considered cost-effective. At the same time, high-priced treatments may preclude NBS from being viewed as cost-effective despite substantial health gains. Strategies to reduce the costs of treatment could impact the value of NBS.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

ACKNOWLEDGEMENTS

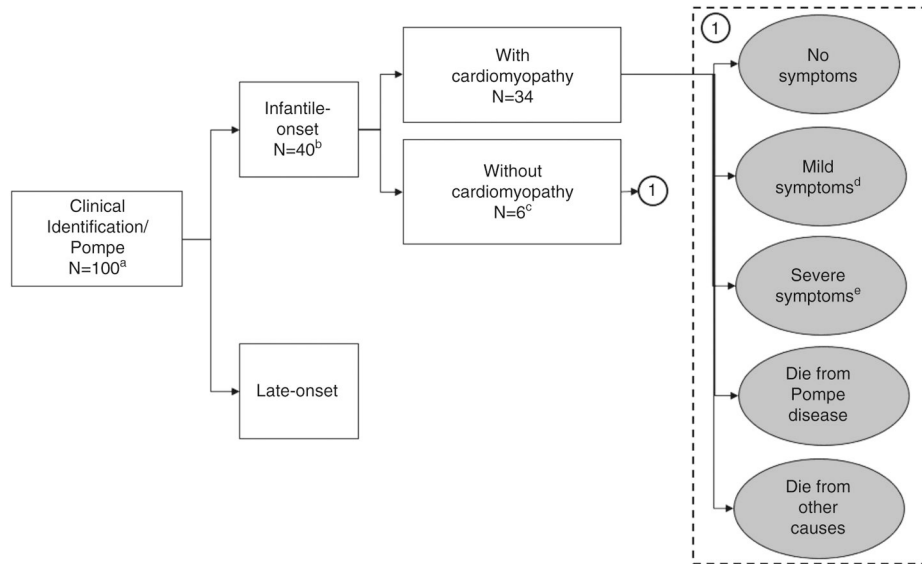
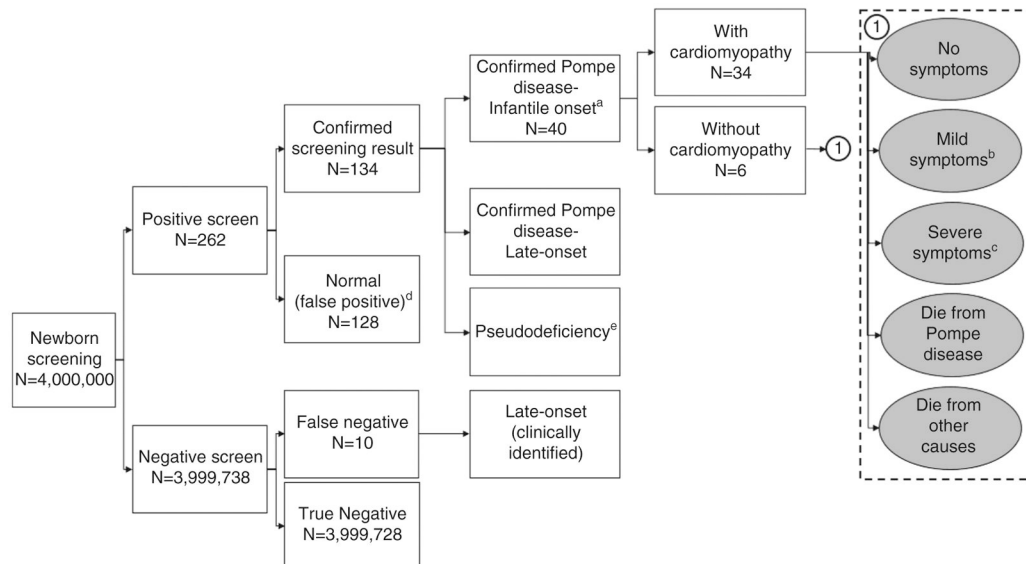
The authors acknowledge the expert panelists for reviewing assumptions and parameter inputs for the simulation model: Priya Kishnani, Deborah Marsden, C. Ronald Scott, Olaf Bodamer, Barry Byrne, Sharon Kardia. We also acknowledge the following individuals for providing information on Pompe disease newborn screening: Janice Bach, (Michigan), Patrick Hopkins (Missouri), Tess Rhodes (Illinois), Sarah Bradley (New York), Suzanne Karabin (New Jersey), and Deeksha Bali (Duke University). We also acknowledge Brittany D'Cruz and Nicole Emblad for their research assistance and Rachel Fisher for her expertise. This work was funded by the Agency for Healthcare Research and Quality (AHRQ), grant number R01 HS020644. The funding agreement ensured the authors' independence in designing the study, interpreting the data, writing, and publishing the report. The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention. The views expressed herein are solely those of the authors and do not necessarily reflect the views of the Secretary of the United States Department of Health and Human Services or of the individual members of the Secretary's Advisory Committee on Heritable Disorders in Newborns and Children.

REFERENCES

1. Berry SA et al. Newborn screening 50 years later: access issues faced by adults with PKU. *Genet. Med* 15, 591–599 (2013). [PubMed: 23470838]
2. NewSTEPs Data Repository. State profiles. <https://data.newsteps.org/newsteps-web/stateProfile/input.action> (2018).
3. US Secretary of Health and Human Services. Charter: Advisory Committee on Heritable Disorders in Newborns and Children. <https://www.hrsa.gov/sites/default/files/hrsa/advisory-committees/heritable-disorders/about/charter.pdf>. (2015).
4. Prosser LA et al. Using decision analysis to support newborn screening policy decisions: a case study for Pompe disease. *MDM Policy Pract.* 3, 2381468318 763814 (2018).
5. US Health and Resources and Services Administration. Newborn screening for Pompe disease: a summary of the evidence and advisory committee decision. <https://www.hrsa.gov/sites/default/files/hrsa/advisory-committees/heritable-disorders/rusp/previous-nominations/pompe-27-june-2018.pdf> (2013).
6. Kemper AR et al. Evidence report: newborn screening for Pompe disease. <https://www.hrsa.gov/sites/default/files/hrsa/advisory-committees/heritable-disorders/rusp/previous-nominations/pompe-external-evidence-review-report-2013.pdf> (2013).
7. Kishnani PS et al. Pompe disease diagnosis and management guideline. *Genet. Med* 8, 267–288 (2006). [PubMed: 16702877]
8. Guo J, Kelton CM & Guo JJ Recent developments, utilization, and spending trends for Pompe disease therapies. *Am Health Drug Benefits* 5, 182–189 (2012). [PubMed: 24991319]
9. Martin JA, Hamilton BE, Osterman MJK, Driscoll AK & Mathews TJ Births: final data for 2015 National Vital Statistics Reports web site. https://www.cdc.gov/nchs/data/nvsr/nvsr66/nvsr66_01.pdf (2017).

10. Neumann P, Gillian S, Russell L, Siegel J & Ganiats T Cost-Effectiveness in Health and Medicine. Vol 2. (Oxford University Press, New York, 2017).
11. Kemper AR, Hwu W-L, Lloyd-Puryear M & Kishnani PS Newborn screening for Pompe disease: synthesis of the evidence and development of screening recommendations. *Pediatrics*. 120, e1327–e1334 (2007). [PubMed: 17974725]
12. Leslie N & Bailey L Pompe disease. University of Washington. GeneReviews. <https://www.ncbi.nlm.nih.gov/pubmed/20301438> (2017).
13. Sullivan W & Payne K The appropriate elicitation of expert opinion in economic models: making expert data fit for purpose. *Pharmacoeconomics*. 29, 455–459 (2011). [PubMed: 21568356]
14. Chiang S-C, Hwu W-L, Lee N-C, Hsu L-W & Chien Y-H Algorithm for Pompe disease newborn screening: results from the Taiwan screening program. *Mol Genet Metab*. 106, 281–286 (2012). [PubMed: 22578805]
15. Kishnani PS et al. A retrospective, multinational, multicenter study on the natural history of infantile-onset Pompe disease. *J Pediatr* 148, 671–676 (2006). [PubMed: 16737883]
16. Chen LR et al. Reversal of cardiac dysfunction after enzyme replacement in patients with infantile-onset Pompe disease. *J Pediatr* 155, 271–5.e272 (2009). [PubMed: 19486996]
17. Wasserstein MP et al. The New York pilot newborn screening program for lysosomal storage diseases: report of the first 65,000 infants. *Genet. Med* 21, 631–640 (2018). [PubMed: 30093709]
18. Kanters TA et al. Burden of illness of Pompe disease in patients only receiving supportive care. *J Inher Metab Dis* 34, 1045–1052 (2011). [PubMed: 21499718]
19. Kishnani PS et al. Timing of diagnosis of patients with Pompe disease: data from the Pompe registry. *Am J Med Genet A* 161, 2431–2443 (2013).
20. Simon NJ et al. Health utilities and parental quality of life effects for three rare conditions tested in newborns. *J Patient Rep Outcomes* 3, 4 (2019). [PubMed: 30671727]
21. Khan AA et al. Higher dosing of alglucosidase alfa improves outcomes in children with Pompe disease: a clinical study and review of the literature. *Genet. Med* 22, 898–907 (2020). [PubMed: 31904026]
22. Sanders GD et al. Recommendations for conduct, methodological practices, and reporting of cost-effectiveness analyses: second panel on cost-effectiveness in health and medicine. *JAMA*. 316, 1093–1103 (2016). [PubMed: 27623463]
23. Weinstein MC & Skinner JA Comparative effectiveness and health care spending—implications for reform. *N Engl J Med* 362, 460 (2010). [PubMed: 20054039]
24. Neumann PJ, Cohen JT & Weinstein MC Updating cost-effectiveness - The curious resilience of the \$50,000-per-QALY threshold. *N Engl J Med* 371, 796–797 (2014). [PubMed: 25162885]
25. Ollendorf DA, Chapman R & Pearson SD Evaluating and valuing drugs for rare conditions: no easy answers. *Value Health* 21, 547–552 (2018). [PubMed: 29753351]
26. National Institute for Health and Clinical Excellence. Interim Process and Methods of the Highly Specialised Technologies Programme updated to reflect 2017 changes. <https://www.nice.org.uk/Media/Default/About/what-we-do/NICE-guidance/NICE-highly-specialised-technologies-guidance/HST-interim-methods-process-guide-may-17.pdf> (2018).
27. Kanters TA et al. Cost-effectiveness of enzyme replacement therapy with alglucosidase alfa in classic-infantile patients with Pompe disease. *Orphanet J Rare Dis* 9, 75 (2014). [PubMed: 24884717]
28. Castro-Jaramillo HE The cost-effectiveness of enzyme replacement therapy (ERT) for the infantile form of Pompe disease: comparing a high-income country's approach (England) to that of a middle-income one (Colombia). *Rev Salud Publica (Bogota)* 14, 143–155 (2012). [PubMed: 23250322]
29. The balancing act of orphan drug pricing. *Lancet* 390, 2606 (2017).
30. Gungor D et al. Quality of life and participation in daily life of adults with Pompe disease receiving enzyme replacement therapy: 10 years of international follow-up. *J Inher Metab Dis* 39, 253–260 (2016). [PubMed: 26531313]
31. Tarnopolsky M et al. Pompe Disease: Diagnosis and management. Evidence-based guidelines from a Canadian expert panel. *Can J Neurol Sci* 43, 472–485 (2016). [PubMed: 27055517]

32. Hermans MM et al. Twenty-two novel mutations in the lysosomal α -glucosidase gene (GAA) underscore the genotype–phenotype correlation in glycogen storage disease type II. *Hum Mutat* 23, 47–56 (2004). [PubMed: 14695532]
33. Karaceper MD et al. The health system impact of false positive newborn screening results for medium-chain acyl-CoA dehydrogenase deficiency: a cohort study. *Orphanet J Rare Dis* 11, 12 (2016). [PubMed: 26841949]
34. Hayeems RZ et al. False-positive newborn screening for cystic fibrosis and health care use. *Pediatrics*. 140, e20170604 (2017). [PubMed: 29025964]
35. Scott CR et al. Identification of infants at risk for developing Fabry, Pompe, or mucopolysaccharidosis-I from newborn blood spots by tandem mass spectrometry. *J Pediatr* 163, 498–503 (2013). [PubMed: 23465405]
36. US Centers for Medicaid and Medicare Services. Physician fee schedule. <http://www.cms.hhs.gov/PhysicianFeeSched/> (2016).
37. IBM Micromedex Redbook. <https://www.ibm.com/products/micromedex-red-book/details> (2017).
38. Kishnani PS et al. Pompe disease diagnosis and management guideline. *Genet. Med* 8, 267–288 (2006). [PubMed: 16702877]
39. US Bureau of Labor Statistics. Current employment statistics. <http://www.bls.gov/we/empsit/cesesummary.htm> (2017).
40. Bach JR, Tran J & Durante S Cost and physician effort analysis of invasive vs. noninvasive respiratory management of Duchenne muscular dystrophy. *Am J Phys Med Rehabil*. 94, 474–482 (2015). [PubMed: 25741618]
41. US Bureau of Labor Statistics. Consumer expenditure survey. <https://www.bls.gov/cex/> (2013).
42. US Centers for Medicare and Medicaid Services. Clinical laboratory fee schedule. <https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/ClinicalLabFeeSched/index.html> (2014).
43. Richardson J, Wittenberg E & Prosser L Health utilities of rare heritable disorders: approaches for health state descriptions and measuring spillover effects on family members. Paper presented at: Society for Medical Decision Making 2016; Vancouver, BC.
44. Prosser LA, Ladapo JA, Rusinak D & Waisbren SE Parental tolerance of false-positive newborn screening results. *Arch Pediatr Adolesc Med*. 162, 870–876 (2008). [PubMed: 18762606]

a Clinical identification**b Newborn screening****Fig. 1. Model framework.**

(a) Clinical identification. ^aCohort of 4,000,000. ^bInfantile-onset Pompe disease was defined as symptom onset prior to 12 months. ^cTwo-thirds of patients identified with infantile-onset Pompe disease without cardiomyopathy experience a delay in treatment and do not start treatment until age 3. ^dMuscle weakness and declining muscle tone, but maintain the ability to walk and perform most activities independently. ^eIncreasing muscle weakness and breathing problems necessitating wheelchair use and ventilator dependence. (b) Newborn screening. ^aInfantile-onset Pompe disease was defined as symptom onset prior to 12 months. ^bMuscle weakness and declining muscle tone, but maintain the ability to walk and perform most activities independently. ^cIncreasing muscle weakness and breathing problems necessitating wheelchair use and ventilator dependence. ^dSince the focus of this study is on

screening for and the identification of infantile-onset Pompe, we have not distinguished the potential reasons for false positives such as being heterozygous for a pathogenic variant or other technical reasons. ^eSix of the ten false negatives are expected to have a pseudodeficiency and never develop any symptoms.

Author Manuscript

Author Manuscript

Author Manuscript

Author Manuscript

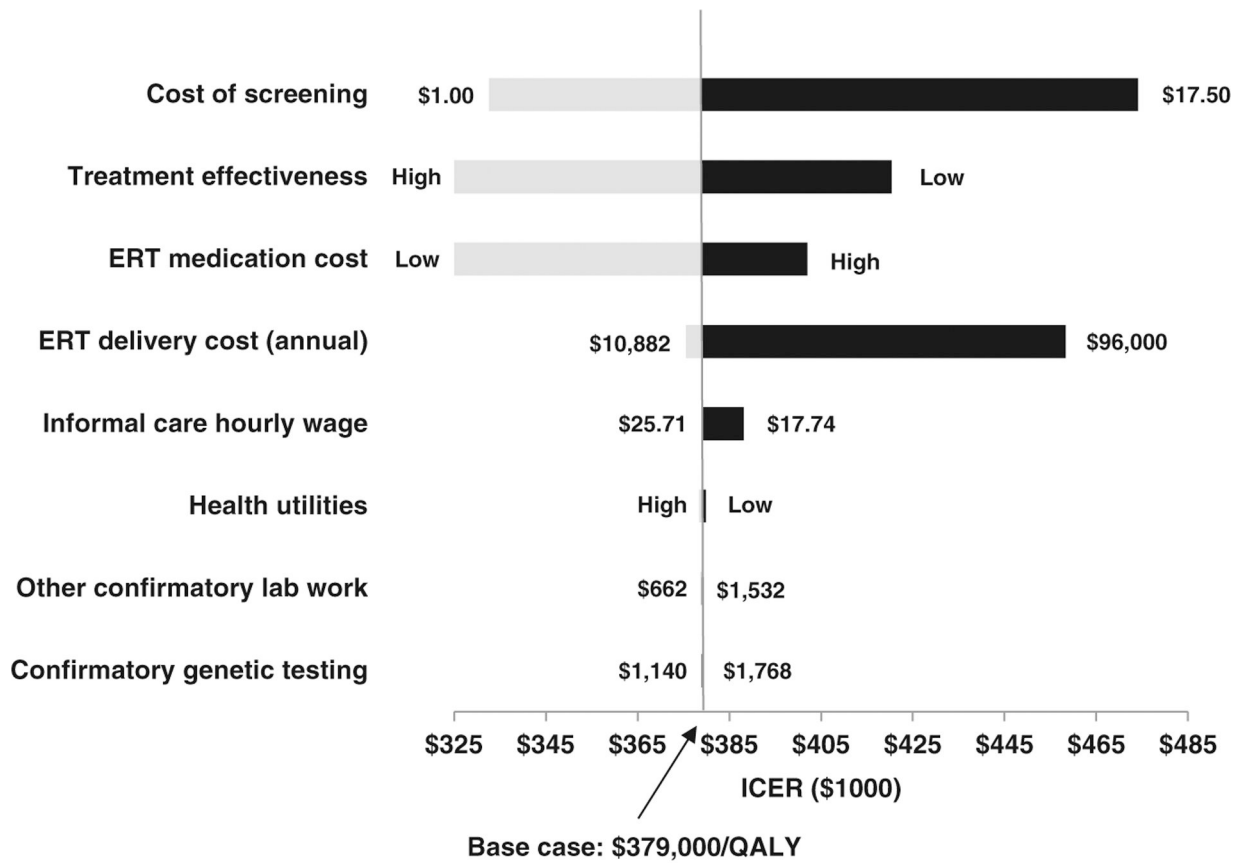


Fig. 2. One-way sensitivity analysis.

ERT enzyme replacement therapy, ICER incremental cost-effectiveness ratio.

Table 1.

Model inputs.

| Parameter | Values | Range | Sources |
|--|---------------|---------------------|-------------------------------------|
| Probabilities^a | | | |
| Incidence of Pompe disease (annual, per 100,000) | 2.5 | 1–2.5 | 35 |
| Screening test characteristics^b | | | |
| Sensitivity | 0.9322 | 0.9315–0.9329 | 14 |
| Specificity | 0.9999 | — | |
| Costs^c | | | |
| Screening test for Pompe disease, per child | \$6.40 | \$6.40–\$17.50 | Personal communication ^d |
| Genetic and other confirmatory tests for newborns who screen positive ^e | \$2,382 | \$1,802–\$3,300 | 36 |
| Enzyme replacement therapy^f | | | |
| Age 1 | \$75,475 | \$71,828–\$117,624 | 37 |
| Age 5 | \$135,856 | \$129,290–\$211,723 | |
| Age 10 | \$256,616 | \$244,214–\$399,922 | |
| Age 15 | \$407,567 | \$387,869–\$635,170 | |
| Age 25 | \$483,043 | \$459,697–\$752,794 | |
| Age 50 | \$513,233 | \$488,428–\$799,843 | |
| Enzyme replacement delivery ^g | \$14,300 | \$10,725–\$96,000 | |
| Pompe disease, mild symptoms | | | |
| Direct medical care (all) ^h | Varies by age | \$7,303–\$14,257 | 38 |
| Informal care, annual ⁱ | \$18,928 | \$12,915–\$18,928 | 18,39 |
| Appointment time, annual ^j | \$2,160 | — | |
| Pompe disease, severe symptoms | | | |
| Direct medical care (all) ^k | Varies by age | \$84,367–\$90,476 | 38 |
| Formal care ^l | \$87,360 | \$57,512–\$87,360 | 40 |
| Informal care ^m | \$151,424 | \$103,318–\$151,424 | 18,39 |

| Parameter | Values | Range | Sources |
|--|---------------|-----------------|---------|
| Medical, non-Pompe disease related | Varies by age | \$1,162–\$6,014 | 41 |
| Watchful waiting ⁿ | \$437 | — | 36,42 |
| Quality adjustments, health utilities^o | | | |
| Mild symptoms with Pompe disease, <18 years | 0.799 | 0.750–0.844 | 43 |
| Mild symptoms with Pompe disease, 18 years | 0.853 | 0.811–0.892 | |
| Severe symptoms with Pompe disease, 0–1 years | 0.399 | 0.341–0.457 | |
| Severe symptoms with Pompe disease, 2–17 years | 0.466 | 0.407–0.525 | |
| Severe symptoms with Pompe disease, 18 years | 0.536 | 0.480–0.594 | |
| QALY loss due to transient positive screen | –0.0005 | — | 44 |

CMP cardiomyopathy, *QALY* quality-adjusted life-year.

^aTransition probabilities for each health state and scenario are provided in Table S1 of the supplemental materials.

^bThe range in sensitivity and specificity corresponds with a range of 0 to 2,800 potential false positives.

^cAll costs were adjusted to 2016 dollars using the Gross Domestic Product Price Deflator.

^dCorrespondence with New Jersey, New York, Michigan, and Missouri newborn screening programs.

^eSee Table S3b for details.

^fAll individuals with confirmed infantile-onset Pompe were assumed to begin receiving 20 mg/kg of alglucosidase alfa every other week at diagnosis (mean = 22 days of life). Redbook wholesale acquisition cost with a 23% discount was used for the base case. A one-time cost of \$7,594 for immune tolerance induction was also applied in the first year of life to 25% of those with infantile-onset Pompe disease and cardiomyopathy. For additional details, see Table S3d.

^gEnzyme replacement therapy (ERT) infusion lasts 6 hours; first year 50–75% at home; after that 100% at home; see supplementary tables for additional details.

^hSee Table S3f for microcosting details. Includes one-time transition costs (e.g., environmental changes and extra medical equipment).

ⁱ14 hours of informal care per week, similar to what is reported by Kanters et al.¹⁸ multiplied by average hourly wage of \$25.71.³⁹

^jBased on average hourly wage of \$25.71.³⁹

^kSee Table S3f for microcosting details. Includes one-time transition costs (gastrostomy, wheelchair equipment, environmental changes).

^l8 hours per day, 7 days per week.

^m16 hours per day, 7 days a week.

Author Manuscript

Author Manuscript

Author Manuscript

Author Manuscript

Additional health-care utilization for patients diagnosed with Pompe disease but without symptoms, annual estimate: one additional outpatient visit; lab work including creatine kinase-MB, comprehensive panel, and urine hex4. There is an additional \$51 cost for appointment time.

⁰In secondary analysis, additional spillover family disutility was also applied to this health state. Spillover values are given in Table S3g.

Table 2.

Projected health and economic outcomes.

| Outcomes | Clinically diagnosed and treated | Newborn screening and treated |
|--|----------------------------------|-------------------------------|
| Health outcomes, infantile-onset | | |
| Screening outcomes (or within first year of life for clinically diagnosed) | | |
| Cases, infantile-onset with cardiomyopathy | 34 | 34 |
| Cases, infantile-onset without cardiomyopathy ^a | 2 | 6 |
| Proportions with cardiomyopathy, by health state, at age 10 | | |
| No symptoms | 0.00 | 0.30 |
| Mild symptoms | 0.19 | 0.67 |
| Severe symptoms | 0.10 | 0.00 |
| Dead from Pompe disease | 0.70 | 0.02 |
| Dead from other causes | 0.01 | 0.01 |
| Health outcomes | | |
| By age 3 | | |
| Ventilator dependent, <i>n</i> | 7.11 | 0.53 |
| Deaths, <i>n</i> | 6.93 | 0.00 |
| By age 10 | | |
| Ventilator dependent, <i>n</i> | 4.87 | 0.17 |
| Deaths, <i>n</i> | 11.58 | 0.00 |
| By age 20 | | |
| Ventilator dependent, <i>n</i> | 4.06 | 0.32 |
| Deaths, <i>n</i> | 14.40 | 0.02 |
| Disaggregated results among individuals diagnosed with infantile-onset Pompe disease (per person)^b | | |
| Screening and confirmation costs | NA | \$2,388 |
| Enzyme replacement therapy (ERT) | \$6,462,230 | \$10,985,209 |
| Medical, Pompe disease other than ERT | \$385,039 | \$178,834 |
| Medical, non-Pompe disease | \$40,731 | \$72,952 |
| Formal caregiving | \$283,564 | \$26,534 |
| Informal caregiving | \$774,734 | \$428,162 |
| Nonmedical (e.g., appointment time costs and environmental modifications) | \$33,318 | \$44,607 |

| Outcomes | Clinically diagnosed and treated | Newborn screening and treated |
|---|----------------------------------|-------------------------------|
| Total costs, health-care sector ^c | \$7,171,563 | \$11,265,916 |
| Total costs, societal | \$7,979,615 | \$11,738,685 |
| Mean QALYs | 15.01 | 26.67 |
| Infantile-onset with cardiomyopathy | 13.76 | 26.65 |
| Infantile-onset without cardiomyopathy | 22.11 | 26.79 |
| Summary and disaggregated results for annual US newborn cohort (n = 4,000,000)^e | | |
| Costs | | |
| Screening and confirmation tests | \$0 | \$26,224,278 |
| Enzyme replacement therapy | \$258,489,185 | \$439,408,352 |
| Pompe disease-related medical care | \$15,747,528 | \$7,639,698 |
| Non-Pompe disease-related medical care | \$291,659,614,973 | \$291,660,903,800 |
| Formal caregiving costs | \$11,342,569 | \$1,061,343 |
| Informal caregiving | \$30,989,365 | \$17,126,462 |
| Nonmedical (e.g., appointment time costs and environmental modifications) | \$1,875,353 | \$2,355,840 |
| Total costs (health-care sector) ^f | \$291,945,194,255 | \$292,135,237,471 |
| Total costs (societal) | \$291,978,058,973 | \$292,154,719,773 |
| Incremental costs (health-care sector) | — | \$190,043,216 |
| Incremental costs (societal) | — | \$176,660,800 |
| Total QALYs | 121,806,506 | 121,806,973 |
| Incremental QALYs | — | 466 |
| ICER ^g (health-care sector) | — | \$408,000/QALY |
| ICER ^g (societal) | — | \$379,000/QALY |

ICER incremental cost-effectiveness ratio, QALY quality-adjusted life-year.

^aUnder clinical identification, 4 of the 6 cases are anticipated to remain undiagnosed until age 3.

^bAverage per person costs and QALYs. Costs and QALYs discounted at 3%.

^cExcludes informal caregiving and nonmedical costs.

^dWeighted average of two cases treated near birth and four cases treated at age 3.

^eCosts and QALYs are discounted at 3% to 2016 USD.

The health-care sector perspective includes all but nonmedical and informal caregiving costs.
ICERs were calculated using additional decimal places for incremental QALYs—see Supplementary tables for detailed results.

Author Manuscript

Author Manuscript

Author Manuscript

Author Manuscript