Health Utilities for Children and Adults With Type 1 Diabetes

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Objective: We studied health utilities in patients with type 1 diabetes to understand potential differences in health utilities as function of age, type of respondent (self report vs. proxy report), and method of assessment (direct vs. indirect).

Research Design and Methods: We elicited self-reported health utilities for adults (n = 213) and children (n = 238) with type 1 diabetes, and by parent proxy report (n = 223) for overall quality of life [Health Utilities Index (HUI) Mark 3 and experienced time-trade-off (TTO) questions] and hypothetical complication states (TTO questions).

Results: Mean health utilities for overall quality of life (QOL) ranged from 0.81 to 0.91. Children had significantly higher overall QOL compared with adults (0.89 vs. 0.85, P<0.01) by HUI, but had no significant difference in QOL by TTO. There were no significant differences in QOL between child self report and parent proxy report. Utilities were higher for HUI versus TTO for parent proxy report (P<0.01) but not for adult or child self report. Utilities for hypothetical complication states were lower than for current QOL. Values were lower for stroke (0.34 to 0.53), end stage renal disease (0.47 to 0.55), and blindness (0.52 to 0.69) than for amputation (0.73 to 0.82) and angina (0.74 to 0.80). Complication utilities for parent proxy report were higher compared with adult self report for most hypothetical complication states.

Conclusions: Individuals with type 1 diabetes with few complications report a relatively high QOL; however, future end stage complications are rated as having a significant impact on QOL. Differences in utilities by age, self report versus proxy report, and method raise important questions about whose utilities should be used in economic analyses.

Key Words: diabetes, health utilities, quality of life

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on a smaller number of respondents, or did not elicit specific utilities for diabetes complication states.

There is a paucity of health utility data from children and adults with type 1 diabetes. As a majority of individuals develop the disease during childhood, a large number of interventions are being conducted solely in pediatric populations, or in populations consisting of children and adults, thus posing multiple dilemmas for investigators conducting cost-effectiveness analyses. Standardized methods for assessing health utilities have been developed primarily for adult populations, but it is unclear whether adult utilities should be applied to children as well. As type 1 diabetes may impact QOL differently for children versus adults, it is critical to understand how health utilities may differ between the 2 groups. Furthermore, because proxy respondents are often used to measure health utilities for younger children, it is important to understand whether health utilities might differ between child self report and proxy report. We therefore had 2 overarching goals for this study: (1) to provide health utilities specific to individuals with type 1 diabetes both for overall QOL and for complication states; and (2) to use type 1 diabetes as a disease paradigm for understanding potential differences in health utilities as function of age, type of respondent (self report vs. proxy report), and method of assessment (direct vs. indirect).

We hypothesized that children with type 1 diabetes would report higher health utilities than adults, that health utilities generated by proxy report would be similar to those generated by child self report, and that direct versus indirect methods would yield similar health utility estimates.

**RESEARCH DESIGN AND METHODS**

We conducted face-to-face interviews with individuals (>8 y) with type 1 diabetes and their parents/guardians (proxy report) who enrolled in the Juvenile Diabetes Research Foundation Continuous Glucose Monitoring trial. Further details of the trial have been described in previous publications, and utilities for the treatment arm have been used for a cost-effectiveness analysis of the trial.

At baseline, indirect and direct methods were used to assess health utilities for overall QOL with diabetes, and direct methods were used to assess health utilities for hypothetical diabetes complication states. Utilities for control and intervention cohorts were combined for this analysis.

**Indirect Measurement**

For indirect measurement of health utilities, all subjects and parents of children 8 to 18 years completed the Health Utilities Index (HUI) Mark 3, an 8 item self-administered questionnaire which assesses health related QOL. Levels of functioning are measured across a variety of attributes, including vision, hearing, speech, ambulation, dexterity, emotion, cognition, and pain, and responses are transformed into a utility score.

**Direct Measurement**

The Time-Trade-Off (TTO) method was selected over other methods because of its unique balance of relative understandability, sensitivity to preference, and use in prior preference studies of patients with diabetes. Individuals were read a description of a specific health state and then were asked to give their preference for years of life with that health state compared with a shorter period of time in perfect health. The response frame for the TTO exercise was linked to life expectancy for different age groups. Younger children had a longer time frame (eg, 40 y) to consider than older adults (eg, 20 y), but the iterative approach to finding the equilibrium point was identical. Individuals aged 8 to 15 years were asked their preference for living 50 years with the health state versus living 50 years in perfect health. A time frame of 40 years was used for individuals 16 to 24 years, a time frame of 30 years was used for individuals 25 to 35 years, and a time frame of 20 years was used for individuals 35 and older. A ping-pong method was used to arrive at the point of indifference where time in the current health state and decreased time in perfect health were equally desirable; this point was then used to calculate the utility score (eg, if 25 y of life in perfect health equals 50 y with diabetes, the utility would be 0.50).

To evaluate overall QOL (experienced TTO), individuals were asked to think about their current health with diabetes. To evaluate QOL associated with diabetes complications (complication TTO), standardized hypothetical state descriptions were used. Hypothetical states have been repeatedly used in TTO studies related to diabetes treatments and complications, allowing for an evaluation of patients’ perceptions of complication states, particularly in populations with a low rate of complications. The functional and symptomatic experience of living with specific diabetes complications (blindness, end-stage renal disease [ESRD], chronic angina/myocardial infarction, stroke, and lower extremity amputation) was described with no specific age attached to the scenario. Complication health utilities were elicited for adult self report, but as well for parent proxy report. Parents must serve as proxy decision makers for their children when considering enrollment in interventional studies that may prevent complications, underscoring the need for assessing health utilities by proxy report. Experienced and complication utility data were collected from all subjects and parents with the following exceptions: (1) subjects <15 years of age were excluded from the experienced TTO questionnaire, as it was unclear whether they would be able to cognitively complete the questionnaires; and (2) subjects <19 years of age were excluded from the hypothetical complication scenario TTO questionnaires due to concerns about the sensitive nature of the health state descriptions.

**Statistical Analysis**

All analyses were performed using Stata 10. To describe the distribution of utilities, mean, median, standard deviation (SD), and interquartile range were calculated. We compared health utilities: (1) for adults versus children; (2) for parent proxy report versus child self report; and (3) for direct (TTO) versus indirect methods (HUI). We also compared parent proxy report versus adult self report for complication states only. Rank sum tests were used to
TABLE 1. Demographics and Clinical Characteristics of the Study Population

<table>
<thead>
<tr>
<th></th>
<th>Children (n = 238)</th>
<th>Adults (n = 213)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (y)</td>
<td>13.7 ± 3.1 y</td>
<td>38.8 ± 13.6 y</td>
</tr>
<tr>
<td>Female [% (n)]</td>
<td>51.7% (n = 123)</td>
<td>58.7% (n = 125)</td>
</tr>
<tr>
<td>Race [% (n)]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>92.4% (n = 220)</td>
<td>96.2% (n = 205)</td>
</tr>
<tr>
<td>Black</td>
<td>1.7% (n = 4)</td>
<td>1.4% (n = 3)</td>
</tr>
<tr>
<td>Asian</td>
<td>0.8% (n = 2)</td>
<td>0.5% (n = 1)</td>
</tr>
<tr>
<td>Other</td>
<td>5% (n = 12)</td>
<td>1.9% (n = 4)</td>
</tr>
<tr>
<td>Mean duration (y)</td>
<td>6.4 ± 3.4</td>
<td>21.6 ± 12.5</td>
</tr>
<tr>
<td>Complications [% (n)]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>0.4% (n = 1)</td>
<td>18.3% (n = 39)</td>
</tr>
<tr>
<td>Hypercholesterolemia</td>
<td>0.8% (n = 2)</td>
<td>32.4% (n = 69)</td>
</tr>
<tr>
<td>Cardiovascular Disease</td>
<td>0.4% (n = 1)</td>
<td>7.0% (n = 15)</td>
</tr>
<tr>
<td>Renal disease</td>
<td>1.3% (n = 3)</td>
<td>8.0% (n = 17)</td>
</tr>
<tr>
<td>Neurological disease</td>
<td>1.7% (n = 4)</td>
<td>3.8% (n = 8)</td>
</tr>
<tr>
<td>Retinopathy</td>
<td>0%</td>
<td>6.1% (n = 13)</td>
</tr>
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</table>

comparing utilities for independent samples and sign rank tests were used to compare paired data (ie, parent proxy vs. child self report). We also performed Spearman correlations of health utilities for parent proxy report versus child self report using the HUI, and for HUI versus experienced TTO. Finally, we assessed the reliability of the TTO and HUI for the control group at baseline and 6 months later. Because of multiple comparisons, significance was defined as a P value < 0.01 for all comparisons.

RESULTS

Table 1 describes the sample and their demographic/clinical characteristics. Mean baseline HbA1c was 7.5 ± 0.9%, reflecting the relatively good control of the cohort overall. As expected, duration of diabetes was longer and rates of complications were higher for adults compared with children. Overall, 12% (n = 52) of subjects had at least 1 major complication.

All adults completed the health utilities questionnaires for overall QOL, and the majority (93%) of hypothetical complication state questionnaires. Most children from 8 to 18 years of age completed the HUI (97%, n = 231); a smaller proportion of children from 15 to 18 years of age completed the experienced TTO (82%, n = 95). Subjects with missing data tended to be younger, with mean ages of 9.7 years (n = 7) for the HUI and 14.5 years (n = 21) for the experienced TTO. The majority of parents completed the HUI (n = 223) and TTO (n = 221) questionnaires; a smaller proportion (74% to 78%) completed the complication state questionnaires.

Utilities for Experienced Overall QOL

Health utility scores for overall QOL for adult self report (≥ 19 y), parent proxy report (8 to 18 y), and child self report (8 to 18 y) are shown in Figure 1 and Table 2. Health utilities as measured by the HUI were relatively high in all 3 groups. For overall current QOL, anywhere from 14% to 39% of individuals reported the maximum health utility (ie, perfect health).

Comparisons by Age

Compared with adults, children had significantly higher utilities (0.89 vs. 0.85, P < 0.01) using the HUI. There were no significant differences using the TTO.

Comparisons for Self Report Versus Proxy Report

There were no significant differences between child self report and parent proxy report using either HUI or TTO. Correlations between child self report and parent proxy report were 0.34 (95% CI: 0.22 to 0.45) for HUI and 0.31 (95% CI: 0.10 to 0.50) for experienced TTO, with scatter plots demonstrating considerable variability between parent and child (Figs. 2A, B). For the HUI, parent-proxy utilities were higher than self-reported child utilities in 41% of cases, equal in 28% of cases, and lower in 31% of cases. For experienced TTO, parent-proxy utilities were higher than self-reported child utilities in 45% of cases, equal in 20% of cases, and lower in 30% of cases.

FIGURE 1. A and B, Boxplots of health utilities for overall quality of life for adult subjects by self report (≥ 19 y), pediatric subjects by parent proxy report (8 to 18 y), and pediatric subjects by self report (8 to 18 y).
Comparisons for HUI Versus TTO

We did find that utilities were higher for HUI versus TTO for parent-proxy report (P < 0.01), but not for child or adult self report. Accordingly, correlations between HUI and TTO were higher for parent-proxy report (0.12 (95% CI: 0.03-0.29)) and child self report (0.20 (95% CI: 0.10-0.39)) than for parent proxy report 0.12 (95% CI: 0 to 0.25). Again there was considerable individual variability (Figs. 3A–C). Self-reported utilities for HUI were higher than TTO in 51% of cases but were lower in 49% of cases, and parent proxy-reported utilities for HUI were higher than TTO in 59% of cases and lower in 41% of cases.

We also assessed test-retest reliability comparing baseline and 6-month values (control group only) for experienced TTO and HUI. Correlations for experienced TTO by self report and proxy report were 0.67 (95% CI: 0.47-0.81) and 0.49 (95% CI: 0.33-0.63), respectively. Correlations for HUI by self report and proxy report were 0.42 (95% CI: 0.25-0.57) and 0.49 (95% CI: 0.33-0.63), respectively.

Utilities for Hypothetical Diabetes Complications

Compared with health utilities for overall QOL, health utilities for hypothetical complication states were lower (Table 1). The complication considered the most damaging to QOL was stroke, while the least damaging complications were angina and amputation. The ranking of complication utilities were similar for adult subjects and parent proxies. The percentage of individuals with health utilities at the maximal value was generally lower for the hypothetical health states versus overall QOL.

Subgroup Comparisons of Utilities for Hypothetical Diabetes Complication States

For the hypothetical complication states, utilities were more favorably rated by proxy report than by adult self report across most states (P < 0.001 for blindness and stroke, P < 0.01 for end stage renal disease, P = 0.03 for angina, and P < 0.01 for amputation).

TABLE 2. Health Utilities for Adult Subjects by Self Report (≥ 19 y), Child Subjects by Parent Proxy Report (8 to 18 y), and Child Subjects by Self Report (8 to 18 y)

<table>
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<tbody>
<tr>
<td></td>
<td>Mean (SD)</td>
<td>Median (25th%-75th%)</td>
<td>Percentage at Maximum Value</td>
</tr>
<tr>
<td>Current overall quality of life</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HUI</td>
<td>0.85 (0.17)*</td>
<td>0.91 (0.79-0.97)</td>
<td>14 (30)</td>
</tr>
<tr>
<td>Experienced TTO</td>
<td>0.81 (0.25)</td>
<td>0.92 (0.73-0.98)</td>
<td>33 (70)</td>
</tr>
<tr>
<td>Hypothetical complication states</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blindness</td>
<td>0.52 (0.33)‡</td>
<td>0.50 (0.23-0.83)</td>
<td>22 (40)</td>
</tr>
<tr>
<td>End stage renal disease</td>
<td>0.47 (0.32)‡</td>
<td>0.48 (0.23-0.73)</td>
<td>8 (16)</td>
</tr>
<tr>
<td>Angina</td>
<td>0.74 (0.28)</td>
<td>0.85 (0.61-0.98)</td>
<td>23 (48)</td>
</tr>
<tr>
<td>Stroke</td>
<td>0.34 (0.31)‡</td>
<td>0.23 (0.03-0.56)</td>
<td>4 (9)</td>
</tr>
<tr>
<td>Amputation</td>
<td>0.73 (0.30)‡</td>
<td>0.88 (0.50-0.98)</td>
<td>27 (57)</td>
</tr>
</tbody>
</table>

*Statistically significant differences in utilities comparing child self report with adult self report (P < 0.01).
†Statistically significant differences in utilities comparing general quality of life comparing the HUI with the TTO (P < 0.01).
‡Statistically significant differences in utilities for hypothetical complication states comparing parent proxy report with adult self report for blindness and stroke (P < 0.001), end stage renal disease (P < 0.01), and amputation (P < 0.01).

HUI indicates Health Utilities Index; TTO, time-trade-off.

FIGURE 2. A and B, Scatter plots of parent proxy-report health utilities versus child self-report health utilities (for overall quality of life) using the Health Utilities Index (8 to 18 y) (n = 216) or Experienced Time-Trade-Off (15 to 18 y) (n = 80).
DISCUSSION

This is one of the first studies to provide empirically-derived health utility data for overall QOL and diabetes-specific complications from adults and children with type 1 diabetes and their parents as proxy respondents. We found that mean health utilities for overall QOL for individuals in our study were relatively high and were comparable to 2 different smaller studies of the United States and Canadian adults with type 1 diabetes which also used the TTO methodology for generating utilities. These studies reported overall utilities of 0.88 (n = 72) and 0.87 (n = 85), which were comparable with the utilities from our study. Our study further extends this work with a much larger sample size and a greater variety of respondents, including children and parents as proxy respondents.

Our findings, however, contrast with the findings of additional studies that have elicited health utilities for individuals with type 1 diabetes. Another study of Canadian adults with type 1 diabetes also used the HUI, estimating an overall QOL health utility of 0.78 versus 0.85 for adults in our study, which may be due to higher complication rates in that population. Similarly, Coffey et al estimated utilities using the Self-Administered Quality of Well-Being index (QWB-SA) in a slightly younger population (mean age, 34 y) of individuals with type 1 diabetes, and reported mean health utility scores of 0.64 to 0.67 for individuals without microvascular, neuropathic, or cardiovascular complications. Finally, Wu et al estimated health utilities of 0.73, 0.68, and 0.64 for adult patients with type 1 diabetes aged <45 years, 45 to 64 years, and ≥65 years, respectively, by predicting QWB scores based on responses to the SF-36, a generic health status assessment instrument that profiles 8 health domains for an individual. The characteristics of our population and the differing methodologies used for obtaining health utilities may account for some of the differences.

We note that adult individuals in our study had higher levels of overall QOL compared with the general population, with 1 recent study reporting overall utilities of 0.81 using the HUI3 in a national probability sample of United States adults.

Individuals with type 1 diabetes did report lower health utilities when asked to consider hypothetical scenarios of complication states. Huang et al also estimated health utilities using hypothetical complication states for individuals with type 2 diabetes, and found a similar ranking of complication states as in our study. However, our rankings were different from those reported by Coffey et al (lowest to highest: blindness, amputation, stroke, and end stage renal disease), who used regression techniques rather than TTO to estimate the decrement in QWB-SA scores for individuals suffering from specific complications.

FIGURE 3. A to C, Scatterplots of health utilities using the Health Utilities Index 3 versus Experienced Time-Trade-off (for overall quality of life) for child self report (15 to 18 y) (n = 95), adult self report (n = 213), and parent proxy report (n = 221).
We found that health utilities for child self report (HUI) were higher than for adult self report, which is consistent with the previous literature demonstrating that health utilities tend to be higher for younger versus older individuals. This makes logical sense given that children with type 1 diabetes, particularly in this cohort, were relatively healthy with lower rates of complications compared with older adults. However, this could also be confounded by disease duration, which was longer for adults versus children.

Although a variety of studies have compared QOL assessments for children with type 1 diabetes with those from their parents as proxy, we are unaware of studies that have compared actual health utilities for this specific population. Consistent with our hypotheses, we did not find significant differences in health utilities between child self report and parent proxy report. Other health preference studies of children with chronic diseases, including those with pediatric brain tumors or extremely low-birthweight infants, have reported higher utilities for parent proxy report compared with child self report, whereas other studies have reported lower utilities.

We found that parent-proxy utilities tended to be higher compared with adult self report for hypothetical complication states. Parents were asked to think about their child’s health for the scenario; these higher scores reflect the fact that parents are generally willing to trade very little time from the length of their child’s life. Another possibility is that the longer life expectancy assumed for children compared with adults in the scenarios may have affected their responses. These differences will potentially have implications for future economic analyses of type 1 diabetes. For example, the higher utilities associated with complication states reported by parents would lead to less favorable (higher) absolute estimates of cost-effectiveness for a specific treatment or therapy. Furthermore, the higher utilities could result in smaller estimated benefits of therapies and a ceiling effect whereby improvements in QALYs over the course of an intervention would be underestimated because of higher baseline estimates of QOL.

Consistent with our hypotheses, we found a lack of differences between HUI and TTO for child and adult self report. However, we did find higher utility scores for HUI versus TTO for parent proxy report, which contrasts with the findings of a recent systematic analysis which reported that direct methods tend to result in higher health utilities compared with indirect methods.

That study incorporated results from studies of adults from patient groups and the general public, focused on individuals with diseases other than type 1 diabetes, and evaluated additional instruments besides the HUI, which may account for the differences in our findings. Further studies are needed to understand why these differences exist for parent proxy report.

Despite the fact that we found significant correlations between direct and indirect utilities at the group level, we found poor correlations at the individual level. This discrepancy has been reported by other studies. It has been suggested that this difference is due to the fact that there can be individual variability in a single utility measurement based on TTO, whereas HUI scores remove this individual variability as they are derived from the mean preferences of a large adult community. Another possibility is that HUI scores, which were generated based on an adult population, are not generalizable to older children or proxy respondents.

Our findings raise the critical question of “whose values” and “what methods” should be used for cost-effectiveness analysis (CEA). The formal recommendation for CEA from the societal perspective is that preferences should be assessed for the general population rather than a patient population and through indirect rather than direct methods. However, others have advocated that in cases where patients’ preferences represent an important outcome (ie, randomized trials) preferences should be derived from affected patient populations through direct methods. For pediatric diseases like type 1 diabetes, the answer to this question is even more complicated, as not only the choice of instrument, but also the choice of which respondent (child self report or parent proxy) needs to be considered. Currently, there is no universal or recommended standard regarding which utilities should be used for pediatric CEA.

Owing to differences that we found, we recommend that investigators consider including both sets of utilities in sensitivity analyses. However, it is clear that further research is needed to further develop methods for eliciting health utilities among children and provide standardization of methods for conducting CEA among children.

We do acknowledge limitations of our study. First, because there is no gold standard for measuring preferences, the validity of utility measurements cannot be directly assessed. However, the rank ordering of the complication utilities has face validity and is consistent with the findings of other studies that have used TTO methods. Secondly, the cross-sectional nature of the data is also a limitation. Third, our correlation plots demonstrated substantial variation between utilities derived from the HUI versus TTO and between children and their parents. However, we did find fair test-retest reliability when comparing utilities at baseline and at 6 months for the control group, suggesting that the differences we found may represent true differences rather than an artifact of poor measurement. Fourth, the significant proportion of individuals with health utility values at the maximum value limits the sensitivity of CUAs for detecting improvements in QOL over time related to specific treatments or interventions. Finally, we did not administer the complication scenarios to children.

We recognize that the preferences we elicited may not be representative of all individuals with type 1 diabetes, as individuals in this cohort were participants in a randomized controlled trial, were receiving intensive insulin therapy, and had better than average control. However, our findings are relevant for the growing population of patients with type 1 diabetes who are adopting new diabetes technologies to help control their disease.

Strengths of this study include the use of both direct and indirect elicitation methods, and the inclusion of both adults and children with type 1 diabetes, and parents of
children with type 1 diabetes. Given the differences in health utilities for the hypothetical complication states that we found for parent proxy report and adult self report, further studies are needed to explore the unique preferences of children regarding diabetes complication states.

CONCLUSIONS
Although individuals with type 1 diabetes report a relatively high QOL, complications of diabetes have a significant impact on the QOL. The differences in health utilities that we found for children versus adults, self report versus proxy report for complication states, and for direct and indirect methods raise important questions about whose utilities should be used in economic analyses. The health utility data generated from this study will be critical for future studies assessing the economic value of current and future interventions targeted at individuals with type 1 diabetes for the health-care system. There is no consensus yet as to how to incorporate health values for adults, children, and their caregivers in economic analyses. Further work is needed to explore reasons for these differences and their potential impact on the economic value of various health care interventions for type 1 diabetes.

APPENDIX

The JDRF Continuous Glucose Monitoring Study Group
Clinical Centers: Listed in order of number of patients enrolled with clinical center name, city, and state. Personnel are listed as (PI) for Principal Investigator, (I) for co-Investigator and (C) for Coordinators:

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University of Minnesota Central Laboratory: Michael Steffes, MD, PhD, Jean M. Bucksa, CLS, Maren L. Nowicki, CLS, Carol Van Hale, CLS, Vicky Makky, CLS.

Cost-effectiveness investigators: National Opinion Research Center, University of Chicago: Michael O’Grady, PhD; Elbert S. Huang, MD, MPH; Anirban Basu, PhD; David O. Meltzer, MD, PhD; Priya John, MPP, Aaron Winn, MPP, Kirsten Rhee, BA. University of Michigan: Joyce M. Lee, MD, MPH.

Juvenile Diabetes Research Foundation, Inc.: Aaron J. Kowalski, PhD.

Operations Committee: Lori Laffel, MD, MPH (co-chair), William V. Tamborlane, MD (co-chair), Roy W. Beck, MD, PhD, Aaron J. Kowalski, PhD, Katrina J. Ruedy, MSPH. Data and Safety Monitoring Board: Ruth S. Weinstock, MD, PhD (chair), Barbara J Anderson, PhD; Davida Kruger, MSN, APRN; Lisa LaVange, PhD; Henry Rodríguez, MD.

Health State Descriptions for Overall Diabetes-Related Complications

**Blindness**
- Imagine a life with blindness:
  - You would not be able to read, see the TV, or drive a car
  - You may also need assistance with many day-to-day tasks such as cooking, cleaning, dressing yourself, and bathing
  - You may also need assistance taking your medications

**End-stage Renal Disease**
- Imagine a life with kidney failure:
  - You would experience fatigue, bone problems, joint problems, itching, and “restless legs”
  - You would need to have dialysis 3X/week and this procedure usually lasts 3-5 h. You may need to make changes in your work or home life to maintain this schedule
  - During dialysis you are attached to a machine. To do this, a needle is inserted into tubing that has been placed under the skin of your arm. The machine then filters the blood to get rid of waste products
  - You may sometimes feel sick or tired for a few hours after you have had dialysis

**Angina**
- Imagine living with chest pain related to heart disease:
  - You experience chest pain after walking a block or 2 but can relieve the pain by stopping or taking medicine
  - Your energy level may be low some of the time
  - You can bathe and dress yourself, and feed yourself without difficulty

**Stroke**
- Imagine life after having a severe stroke:
  - You cannot move the arm or leg on the side that you write with
  - You can stand with a leg brace and walk a short distance with help
  - You can use a wheelchair. You cannot climb stairs
  - You need help to dress, bathe, and use the bathroom. You need help preparing and eating food
• You might have difficulty speaking or finding the right words.

Amputation

Imagine a life after you have lost part of your lower leg or foot:

• You may be able to walk with an artificial leg, or you may have to use a wheelchair to get around.

• You might have some difficulty performing daily tasks such as driving, shopping, or cleaning your house.

REFERENCES


