# Title

Modelling diabetes and depression in Pakistan: Using economic modelling to inform intervention design and a clinical trial of a behavioural activation intervention

Author(s):

1. David Glynn, Centre for Health Economics, University of York
2. Pedro Saramago, Centre for Health Economics, University of York
3. Naveed Ahmed, Centre for Health Research and Implementation, Diabetic Association of Bangladesh and BIRDEM General Hospital, Bangladesh
4. Saima Afaq, Department of Health Sciences, University of York and Institute of Public Health and Social Sciences, Khyber Medical University Peshawar Pakistan.
5. Faiza Aslam, Institute of Psychiatry, Rawalpindi Medical University, Pakistan
6. Abdul Basit, Baqai Institute of Diabetology & Endocrinology (BIDE)
7. David Ekers, Department of Health Sciences, University of York and Tees Esk and Wear Valleys NHS FT, United Kingdom
8. Asher Fawwad, Baqai Institute of Diabetology & Endocrinology (BIDE)
9. Naomi Gibbs, Centre for Health Economics, University of York
10. Edward Fottrell, Institute for Global Health, University College London, UK
11. Richard IG Holt, Human Development and Health, Faculty of Medicine, University of Southampton, UK
12. Rowena Jacobs, Centre for Health Economics, University of York
13. Asima Khan, Baqai Institute of Diabetology & Endocrinology (BIDE)
14. Zia Ul-Haq, Institute of Public Health, Khyber Medical University, Pakistan
15. Gerardo Zavala, Centre for Health Economics, University of York
16. Najma Siddiqi, Department of Health Sciences, University of York
17. Simon Walker, Centre for Health Economics, University of York

# Abstract

**Objectives**

The ‘Developing and evaluating an adapted behavioural activation intervention for depression and diabetes in South Asia (DiaDeM)’ trial investigates a psychological intervention, behavioural activation (BA), on people with both diabetes and depression in Bangladesh and Pakistan. This study aimed to aid the intervention and trial design.

**Design**

This was a modelling study using microsimulation to assess the intervention's cost-effectiveness. Diabetes was modelled using the UK Prospective Diabetes Study (UKPDS) model based on Pakistani patients and depression was modelled using Patient Health Questionnaire-9 trajectories allowing for multiple depressive episodes. It was assumed that diabetes-related adverse events increased depression recurrence, while depression impacted HbA1c, increasing diabetes-related events.

The model estimated (i) maximum cost of BA which would be cost-effectiveness (headroom analysis) to inform intervention design, and (ii) value of reducing uncertainty around different measures (value of information analysis) to prioritise data collection in the DiaDeM study

**Setting**

Analysis was conducted from a Pakistani healthcare perspective over a lifetime with costs and outcomes discounted at 3%.

**Interventions**

BA plus usual care was compared against usual care. BA involved six sessions by a trained (non-mental health) facilitator. The usual care comparator was the prevailing mix of pharmacological and non-pharmacological treatments used in Pakistan.

**Primary and Secondary Outcome Measures**

The primary outcome was disability-adjusted life-years (DALYs). Secondary outcomes included life years, healthcare costs and the rate of depression and diabetes-related events.

**Results**

Over their lifetime, individuals receiving BA plus usual care avoid 3.2 (95% credible interval: 2.7, 3.8) years of mild depression and experience fewer diabetes-related events. BA plus usual care resulted in an additional 0.27 (0.03, 0.52) life years, 0.98 (0.45, 1.86) DALYs averted, and had incremental healthcare costs of -$97 (-$517, $142), excluding BA costs. The maximum cost per BA course at which was cost effective is $83 ($9, $214). Value of information analysis found the most important measures to include in the trial are impact of depression on diabetes and PHQ-9 over time.

**Conclusions**

This is the first model to jointly model depression and diabetes for South Asia and uses novel methods to reflect the diseases and inform intervention and trial design. This evidence has helped to inform the design of the DiaDeM intervention and the trial to evaluate it.

# Strengths and limitations:

* The study modelled the relationship between depression and diabetes to better capture the impact of the intervention of interest on both conditions.
* Depression was modelled as a recurrent, episodic condition, enhancing the reliability of results.
* The study captures the health outcomes and costs for individuals with diabetes and depression in Pakistan
* Due to data limitations the analysis was conducted for Pakistan only, however the DiaDeM trial will be carried out in Pakistan and Bangladesh.
* Evidence was synthesised from diverse sources and setting and this raises challenges around the comparability and generalisability to Pakistan

# Main text:

# Introduction

Individuals can experience multiple long term conditions (MLTCs) simultaneously [1, 2]. The co-occurrence of diabetes and depression is recognised as an important issue [3-5]. These conditions can interact in complex ways which can increase morbidity and mortality. MLTCs are increasingly recognised as an important issue globally which requires research [6, 7].

The ‘Developing and evaluating an adapted behavioural activation intervention for depression and diabetes in South Asia’ (DiaDeM) trial is designed to address the co-occurrence of these MLTCs [8]. The DiaDeM trial is taking place in Bangladesh and Pakistan [9]. It aims to compare an adapted behavioural activation (BA) therapy delivered by non-specialist health workers in addition to optimised usual care to optimised usual care alone. BA is a psychological treatment that has been shown to treat depression effectively and can be delivered by non-specialist health workers. BA helps people make the link between what they do and how they feel and supports them to make changes to improve their health [10, 11]. However, evidence is largely from high income countries and may not be generalisable to South Asian low-middle-income countries because of differences in the cultural context (including attitudes towards depression and talking therapies) and healthcare systems. Further, there is limited evidence on BA for the treatment of depression as part of MLTCs, including diabetes [12]. This study will add to the growing literature on cost effectiveness in the context of MLTCs [7, 13, 14].

In this paper we develop a novel decision analytic model and use it to inform the design of the DiaDeM intervention and trial [15]. This model takes account of the natural history of diabetes and depression and any interactions. It predicts costs, morbidity and mortality over a patient’s lifetime, adjusting costs and patient characteristics to match the decision context. There were two aims of the analysis. Firstly, conducting a head room analysis to estimate the maximum cost of a BA treatment which would still be considered cost-effective for a given treatment effect [16, 17]. These helped to inform the number of BA sessions to provide. Secondly, conducting value of information analyses to inform the trial design by estimating the relative value of collecting specific outcome measures during the trial, for example, should we prioritise collecting data on systolic blood pressure or low density lipoprotein [18, 19]. Further, this model will be used to assess the long-term cost-effectiveness of the DiaDeM BA intervention following the culmination of the definitive trial.

# Methods

## Overview

A MLTC diabetes and depression decision analytic model was developed to estimate the lifetime cost and health impacts of a behavioural activation intervention in addition to usual care versus usual care alone in Pakistan for individuals with both diabetes and depression [15]. This analysis focuses on Pakistan only because of lack of data for Bangladesh. Health outcomes included life years and disability-adjusted life-years (DALYs) averted, a generic health outcome capturing morbidity and mortality. Costs reflected those related to health care, including out of pocket payments given the mixed public-private nature of the Pakistan health care system [20]. Costs and outcomes were discounted at a rate of 3% per annum in line with international guidelines [21]

The decision model was based on a previously developed diabetes model and an innovative de novo depression component [14, 22-24]. All patients enter the model with known diabetes and depression. The treatment options modelled are those included in the DiaDeM trial: BA in addition to usual care vs. “optimised” usual care. The BA intervention is delivered in six face-to-face or online sessions by a trained (non-mental health) facilitator. The usual care comparator is the prevailing mix of pharmacological and non-pharmacological treatments used in Pakistan, “optimisation” in this case is the provision of an information leaflet with details for accessing care locally [25].

Next, we describe the diabetes model, then the depression model and finally we describe the nature of their interaction. This model was developed in R and the Viking computing cluster at the University of York was used to carry out all analyses [26].

### Patient and public involvement

The model was produced in collaboration with local researchers and the international DiaDeM advisory group, which included academics, policy makers and patient representatives.

## Diabetes model

The diabetes component of the model is based on the ‘United Kingdom Prospective Diabetes Study’ (UKPDS) Outcomes Model 2 which captures the risk of diabetes complications and mortality over an individual’s lifetime based on their characteristics including general characteristics (e.g. age, gender, years with diabetes) and a range of risk factors and biomarkers (e.g. HbA1c and estimate glomerular filtration rate (eGFR)) [23, 24]. The model captures the risk of the following diabetes complications: congestive heart failure, myocardial infarction, ischaemic heart disease, stroke, blindness, ulcer, amputation and renal failure. In each year patients are at risk of dying and/or having a diabetes related complication. The risk of events depends on patient characteristics and any history of previous events in the model. Risk factors also change over time e.g. eGFR deteriorates with age [24]. To reflect the Pakistani context, patient profiles are based on individuals attending a diabetes clinic in Pakistan (see section on patient population for further details). A similar approach of accounting for national patient characteristics has recently been used to model diabetes in India [27]. To our knowledge this is the first time the UKPDS model has been adapted in this way to Pakistan. A schematic of the diabetes model is shown in appendix Figure A1.

## Depression model

Depression is modelled as a cyclical disease in which patients can potentially experience multiple depressive episodes [14, 28]. Employing a novel approach to depression modelling, we modelled outcomes at the level of individual depressive symptom scores captured by Patient Health Questionnaire-9 (PHQ-9) trajectories over time (see Figure 1). Individuals enter the model in a depressive episode, they then gradually recover (i.e. their PHQ-9 score decreases) with the rate of recovery dependent on the time since the episode began and the treatment they receive. Throughout they are at risk of having a new depressive episode, even if they have not recovered from the previous episode. If they experience a new episode, their PHQ-9 score increases to a value which represents episodic depression for them (this depends on their baseline PHQ-9 score and their lowest PHQ-9 score in the current episode), after which they begin recovering and the cycle starts again. Evidence to inform the initial distribution of PHQ-9, the recovery rate and the risk of a new depressive episode was taken from the INDEPENDENT study, an RCT of individuals with depression and diabetes in India [29] (further details in Appendix A3).

[INSERT FIG1]

## Interaction between diabetes and depression

The model includes a two-way interaction between diabetes and depression. Depression increasing the risk of future diabetes-related events by impacting on an individual’s HbA1c levels, for example, because depression may result in worse self management of diabetes [30]. The occurrence of diabetes adverse events increasing the risk of new depressive episodes. The relationship between PHQ-9 and future HbA1c was based on the INDEPENDENT study [29]. The increased risk of depressive episodes following diabetes-related complications was based on a published study examining the relationship between complications and incidence of depression (see Appendix A3 for further details) [31]. A schematic summarising the mechanism of this interaction is given in Figure 2.

[INSERT FIG2]

### Treatment effect for Behavioural Activation

A large, network meta-analysis (NMA) of psychological therapies found that BA relative to “care as usual” resulted in a standardized mean difference (SMD) of -0.73 with a 95% confidence interval –0.95 to –0.52, where negative values indicate improvement on a continuous outcome scale [32]. To transport this treatment effect into our model we assume that the BA impacts PHQ-9 though modifying the rate of recovery during a depressive episode. A calibration approach was used to “back calculate” the change in rate of recovery with BA required to result in a -0.73 SMD on PHQ-9 [33, 34]. It was found that an SMD for PHQ-9 of 0.95 was consistent with a doubling of the rate of recovery with BA relative to usual care. See Figure 1 for an illustration of the impact of BA on the PHQ-9 time path and see appendix A3 for full details of the analysis, including the characterisation of uncertainty.

## Patient population

Patient profiles were built based on a representative population of individuals with diabetes from Pakistan using the Baqai Institute of Diabetology & Endocrinology (BIDE) patient registry containing 28,942 individuals [35]. Information on disease history and heart rate not captured in the registry were imputed using external data [14, 29, 36]. The BIDE register did not include information on depression, so PHQ-9 scores were estimated for individuals based on the relationship observed between covariates and PHQ-9 in the INDEPENDENT trial [29], with uncertainty in predictions captured. Further, missing covariates were imputed using multiple imputation by chain equations (MICE), which produces imputations accounting for the underlying uncertainty. To account for variability in patients, we randomly sampled patient profiles from the created data set. It was assumed that the joint distribution of patient characteristics in our patient profile data set (based on BIDE and other sources) represented the joint distribution in the population of interest. Full details of the development of the patient population are available in the appendix and descriptive statistics of the patient population for the analysis are provided in Table A2.

## Costs and Quality of life

### Costs

A targeted review of Pakistani costing studies was conducted to identify costs for the model. No studies were identified which provided all the necessary costing information reflecting all of the clinical events associated with diabetes. However, a recent high-quality study looking at general management costs associated with diabetes was identified [37]. This provided an estimate of background diabetes costs (i.e. routine care for diabetes and excluding diabetic events and complications) for Pakistan. Alva et al provided an estimate of background diabetes costs, diabetes event and complication related costs for the United Kingdom [38]. To estimate costs of diabetes related events in Pakistan, we calculated the ratio of background costs for Pakistan (using Gupta et al [37]) and the United Kingdom (using Alva et al [38]) and assumed that the ratio of costs was constant across all events to estimate Pakistan costs. Costs were inflated to 2020 USD values, see appendix for full list of costs (Table A4) and for details on conversion and inflation adjustment.

The cost of care per depressive episode was estimated to be $67.07, this includes inpatient care, outpatient care and costs of medicines [39]. A cost per course of BA of $15 was estimated based on the expected resource use from the pilot trial protocol, which was applied in addition to the cost of usual care in the BA intervention arm for each depressive episode [8].

### Disability weights

Quality of life impacts for each diabetes-related complication were captured in DALY weights [40]. For depression, the PHQ-9 score for each individual in each month was classified into none, mild, moderate and severe. DALY weights were applied to each of these categories and the average DALY score for each year was calculated. In calculating DALYs, the life expectancy for each age group was taken from the World Health Organisation (WHO) global burden of disease 1990-2019 survey to reflect years of life lost [41]. See appendix for full information (Table A5).

## Economic analysis

## Cost-effectiveness analysis

An intervention is considered cost-effective if the health produced by the intervention exceeds the health which could be generated elsewhere using the same resources. Cost-effectiveness was assessed using a cost-effectiveness threshold of $183 per DALY averted [42-44]. This is an estimate of the marginal productivity for the Pakistani health system reflecting how much health could be generated elsewhere if resources were used for alternative purposes i.e. an intervention must avert a DALY for less than $183 or it would not be considered a cost-effective use of resources. Cost-effectiveness is presented in terms of incremental cost-effectiveness ratios (ICER), incremental net health benefits and incremental net monetary benefits. Costs and outcomes were discounted at a rate of 3% per annum [21].

### Headroom analysis

Headroom analysis allows us to estimate the maximum price of a treatment at which it remains cost effective for a given level of effectiveness [16, 17]. Here it is used to estimate the maximum cost at which BA is the cost-effective option, i.e. we estimated the maximum cost of BA for which the ICER of BA versus usual practice is equal to the cost-effectiveness threshold of 183US$. As individuals potentially experience multiple episodes of depression, the total cost is split over the expected number of depressive episodes to calculate a maximum cost for BA per episode. Further details of the headroom analysis are provided in appendix A6.

### Value of information (VOI) analysis

VOI methods quantify the costs of uncertainty in health or monetary terms in terms of the chance and consequences of making a wrong decision (i.e. incorrectly implementing a non-cost-effective treatment). These methods allow for the estimation of the value of collecting information in a trial [18, 19]. Expected value of partial perfect information (EVPPI) methods allow analysts to quantify the value of collecting information on individual outcomes or groups of outcomes. In the case of DiaDeM, we use EVPPI to compare the value of resolving uncertainty in 21 parameter groups (see appendix Table A7.1), to identify those which are potentially most important to collect additional evidence on in the DiaDeM trial. These parameters are classified as short, medium and long term, reflecting the degree to which hypothetical trials with short, medium and long term follow up could gather information on each parameter group. For example, a trial would require a long follow up to provide substantial information on the risk of diabetes events and mortality. However, only a short follow up would be required to observe the costs associated with routine depression care. Note that some aspects of the model (such as the effect of depression on diabetes) are captured by just one parameter, whereas others (such as the evolution of estimated glomerular filtration rate over time) require many parameters.

The VOI analysis is carried out assuming the headroom cost of BA to maximise uncertainty, with alternative costs of BA used for a scenario analyses. To estimate the population EVPPI, we multiply the individual EVPPI estimates by an estimate of the prevalent population with diabetes and depression, this is 12.84 million in Pakistan [45-47]. Full details are provided in appendix A7.

## Generating model predictions

This is a microsimulation model which generates estimates for population outcomes by repeatedly simulating individuals and recording their outcomes. Therefore, it is necessary to check the number of patients required to achieve convergence of population level results. The model was found to have reasonable convergence after simulating approximately 8000 individuals). To reflect uncertainty in model inputs a probabilistic sensitivity analysis (PSA) was carried out by drawing 1000 times from the sampling distribution of each input parameter [15]. Outcomes (e.g. life years, DALYs etc) were computed for each combination of input values resulting in a posterior distribution for each outcome. Point estimates were computed from the mean of these distributions and credible intervals by computing the relevant quantiles.

# Results

## Clinical outcomes

Table 1 summarises the clinical outcomes predicted by the model including depression outcomes, diabetes events and cause of death. As described above, BA is expected to double the rate of recovery from depressive episodes meaning that over their lifetime, those who receive BA are predicted to avoid 3.24 years of mild depression (PHQ-9 > 5) and 0.65 years of moderate depression (PHQ-9 > 10) relative to usual care. BA is expected to reduce the rate of nearly all diabetes events by lowering HbA1c. This reduction in diabetes events increases life expectancy by 0.27 years on average. BA also slightly reduces the rate of depressive episodes though the interaction with diabetes outcomes. Lower PHQ-9 results in lower HbA1c resulting in a lower risk of diabetes events and ultimately a lower risk of a depressive episode.

Table 1: Clinical outcomes simulated in the DiaDeM model, covering depression outcomes, diabetes related outcomes, mortality and cause of death. UC = Usual Care, BA = behavioural activation, PHQ-9 = Patient Health Questionnaire-9, MI = myocardial infarction, IHD = ischaemic heart disease, CHF = congestive heart failure, CrI = credible interval.

|  |  |  |  |
| --- | --- | --- | --- |
| **Outcome** | **UC** | **UC plus BA** | **UC plus BA vs UC (incremental analysis)** |
| **Mean (80% CrI)** | **Mean (80% CrI)** | **Mean (80% CrI)** |
| **Depression outcomes** |  |  |  |
| Years PHQ-9 > 5  (mild depression) | 4.39 (3.69,5.07) | 1.14 (0.91,1.4) | -3.24 (-3.83,-2.7) |
| Years PHQ-9 > 10  (moderate depression) | 0.86 (0.72,1.02) | 0.21 (0.18,0.24) | -0.65 (-0.81,-0.51) |
| Number of depressive events per 1000 person years | 236 (212,264) | 234 (210,261) | -2 (-4,0) |
| **Diabetes events per 1000 person years** |  |  |  |
| First MI | 24 (11.9,36.8) | 23 (11.4,35.4) | -0.9 (-1.8,-0.2) |
| Second MI | 5.3 (1.7,10.1) | 5.2 (1.7,9.9) | -0.2 (-0.5,0.2) |
| First stroke | 15.5 (3.8,31.5) | 14.7 (3.7,29.9) | -0.8 (-1.7,-0.1) |
| Second stroke | 5.4 (0.1,12.4) | 5.2 (0.1,12.4) | -0.2 (-0.7,0.1) |
| CHF | 10.1 (2,22.1) | 10.1 (2,22.2) | 0 (-0.4,0.4) |
| IHD | 10.3 (3.8,18.9) | 10.3 (3.7,18.9) | 0 (-0.4,0.3) |
| First amputation | 11 (1.9,25.9) | 10 (1.8,23) | -1 (-2.5,-0.1) |
| Second amputation | 4.6 (0.6,11.4) | 4 (0.5,10) | -0.6 (-1.5,0) |
| Blindness | 5.9 (1.1,12.3) | 5.4 (1,11.4) | -0.5 (-1.1,0) |
| Renal failure | 4.3 (0.1,12.1) | 4.3 (0.2,11.9) | 0 (-0.2,0.2) |
| Ulcer | 3.8 (0.2,10) | 3.4 (0.2,8.9) | -0.4 (-1.1,0) |
| Cataract | 24.9 (21.5,28.8) | 24.7 (21.4,28.4) | -0.3 (-0.7,0.1) |
| Severe hypo | 14 (13.7,14.3) | 14.1 (13.7,14.4) | 0.1 (-0.4,0.6) |
| **Mortality** |  |  |  |
| Life years | 19.07 (14.38, 23.81) | 19.34 (14.73 23.87) | 0.27 (0.03, 0.52) |

## Cost effectiveness analysis

Table 2 summarises the discounted results comparing usual care against usual care plus BA including and excluding the $15 BA treatment costs. BA plus usual care results in lower overall health care costs and more DALYs averted than usual care. At $15 per course, BA dominates usual care, being less costly and more effective.

Table 2: Summary of cost-effectiveness results comparing BA in additional to usual care versus usual care alone. BA = behavioural activation, ICER = incremental cost effectiveness ratio, CrI = credible interval.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Treatment option** | **Total Healthcare Costs (95% CrI)** | **Incremental costs, over usual care (95% CrI)** | **DALYs averted (95% CrI)** | **Incremental DALYs averted, over usual care (95% CrI)** | **Incremental net monetary benefit, over usual care (95% CrI)** | **ICER** |
| Usual care | $10,839  ($7,205 to $17,088) | - | -3.5  (7.96 to -9.58) | - | - |  |
| BA (excluding BA treatment costs) | $10,742  ($7,187 to $16,803) | -$97  (-$517 to $142) | -2.52  (9.07 to -8.70) | 0.98  (1.86 to 0.45) | $276.63  ($23.92 to $717.12) | Dominates usual care |
| BA (including BA treatment costs of $15) | $10,792  ($7,237 to $16,854) | -$47  (-$467 to $192) | -2.52  (9.07 to -8.70) | 0.98  (1.86 to 0.45) | $226.48  (-$26.22 to $666.97) | Dominates usual care |

## Headroom analysis

As shown in Table 2 the incremental net monetary benefit (INMB) for BA excluding treatment costs was $276.63 ($23.92 to $717.12). The average number of depressive episodes per individual in the BA arm discounted to present value was 3.34 (2.56 to 4.23) resulting in a headroom cost of$82.58 for a course of BA ($8.60 to $214.10). This is the maximum cost per course of treatment per person which would be expected to be cost-effective. It should be noted that this is considerably higher than the estimated $15 based on expected resource use from the DiaDeM pilot trial. Due to uncertainties in the evidence, the headroom estimate was associated with considerable uncertainty with the 95% credible interval ranging from $8.60 to $214.10. This headroom estimate was used to inform the maximum number of sessions per person for the DiaDeM trial.

## Value of information (VOI) analysis

For the base case VOI is calculated based on the head room cost for the intervention $82.58. Table 3 presents results scaled to reflect the prevalent population in Pakistan (12.84 million). EVPPI estimates the value of resolving uncertainty in a group of parameters. All else being equal (e.g. evidence will cost the same to produce, will resolve the same amount of uncertainty etc.) higher EVPPI values for a given group of parameters indicate that there is potentially more value in gathering data on this group. The largest EVPPI value is for risk of diabetes events and mortality ($728.95m). Data on the risk of these events would require long term follow up and is likely to be best collected using a registry rather than as part of a randomised controlled trial. Data could be captured on PHQ-9 time path over a shorter period (say 1 year) and is expected to provide significant value ($473.3m). There is also value in learning the BA treatment effect ($18.16m), which should be feasible over the time period of a trial. There is expected to be significant value in understanding the trajectories of many of the parameters used in the model, the highest being for HDL Cholesterol ($386m). These parameter groups require medium to long term follow up to capture fully. However, shorter trial designs may provide partial information on these outcomes capturing the trajectories over the trial period.

Appendix A7 presents results for four further sensitivity analyses exploring the impact of different assumptions about the cost of BA: 1) $15 based on the expected resource use from the trial intervention, 2) $65.65 found in an Indian study with intensive compliance efforts [48], 3) $8.60 and 4) $214.10, the lower and upper credible intervals from the headroom analysis respectively. In each case the value of further research is lower than the base case, this is because the headroom analysis chooses the BA price which maximises uncertainty.

Table 3: EVPPI results for estimated protocol cost of BA and headroom cost. EVPPI = Expected value of partial perfect information; USD = United States Dollar; BA = Behavioural activation; PHQ-9 = Patient Health Questionnaire-9; HbA1c = Hemoglobin A1c; BMI = Body mass index; LDL = Low density lipoprotein; HDL = High density lipoprotein; eGFR = estimated glomerular filtration rate.

|  |  |
| --- | --- |
| **Group of parameters** | **EVPPI for population in millions USD [rank]** |
| **Headroom cost, $82.58** | |
| Short term parameters |  | |
| BA treatment effect | $18.16m [15] | |
| PHQ-9 time path with usual care | $473.3m [2] | |
| Effect of depression on diabetes | $395.58m [3] | |
| Costs of routine depression care | $0.03m [19] | |
| Costs of routine diabetes care | $0m | |
| Medium term parameters |  | |
| Time path for HbA1c | $243.81m [10] | |
| Time path for BMI | $231.96m [11] | |
| Time path for LDL cholesterol | $354.58m [6] | |
| Time path for systolic blood pressure | $277.29m [9] | |
| Time path for HDL cholesterol | $386m [4] | |
| Time path for haemoglobin | $0m | |
| Time path for white blood cell count | $355.91m [5] | |
| Time path for heart rate | $169.83m [12] | |
| Time path for smoking | $154.04m [13] | |
| Time path for peripheral vascular disease | $37.12m [14] | |
| Time path for microalbuminuria | $321.5m [7] | |
| Time path for atrial fibrillation | $0.81m [18] | |
| Time path for eGFR | $293.04m [8] | |
| Long term parameters |  | |
| Effect of diabetes complications on depression | $6.68m [16] | |
| Costs associated with diabetes events | $5.69m [17] | |
| Risk of diabetes events and mortality | $728.95m [1] | |
| Probability of BA being cost effective | 42% | |

# Discussion

In this paper we developed a novel MLTC model of diabetes and depression to assess the potential benefits of a BA intervention and to carry out a headroom and value of information analyses with the aim of informing the DiaDeM intervention design and trial. The model developed for this paper will also be used to assess the long-term cost-effectiveness of the DiaDeM BA intervention following the culmination of the definitive trial.

From the analysis, BA is expected to result in considerably less time spent in depressive episodes (3.2 years of mild depression and 0.65 years of moderate depression avoided) and is also expected to reduce the occurrence of diabetes-related adverse events. BA was found to improve health outcomes and reduce costs. In the headroom analysis we found that the maximum price at which BA was expected to be cost effective was $82.58 per course of treatment per person based on a treatment effectiveness which doubles the rate of recovery from depression (95% credible interval ranging from $8.60 to $214.10). Typical wage rates for the relevant staff in Pakistan are approximately $2/hour and each session (other than the first session) is expected to take 30 minutes, resulting in a cost of $1 per session (excluding preparation costs). The lower bound estimate for the headroom analysis is $8.60, with a cost per session of $1. A conservative analysis implies that the intervention is expected to be cost effective even if requiring over 8 sessions per person ($8.6/$1 > 8). The VOI analysis found that there was considerable value in collecting additional information on the different parameters in the model including large value in short term outcomes such as the effect of depression on diabetes ($395.58m) and PHQ-9 time path ($473.3m). These assessments fed into both the intervention and trial design for DiaDeM. This analysis also suggests that there is considerable value in reducing uncertainty about the time path of biomarkers such as HbA1c and BMI. This can provide an estimate of the value of setting up longitudinal data collection for each of these measures. This could then be compared against the expected value of other research projects competing for funding [49].

This paper employed a novel approach to predicting outcomes in depression by modelling individual PHQ-9 scores over time. This can be compared to models which consider a single episode of depression [50-52] and models which are based on discrete states of depression [14, 28]. This approach may be more intuitive and by not classifying patients into broader categories we do not lose information and can better model individual level variation in outcomes. This model structure also allowed us to use individual patient data to estimate the impact that changes in depressive symptoms (PHQ-9) have on diabetes (through HbA1c). External data were used to inform the impact of diabetes-related complications on rates of new depressive episodes [31]. This approach to modelling two-way interactions uses separate data sources to estimate the independent effect of diabetes on depression and vice versa. The endogeneity between the MLTCs is then imposed by the model structure. This makes strong implicit assumptions about the underlying causal structure of disease interactions. A more sophisticated approach would require long term longitudinal data on diabetes and depression outcomes. Careful application of causal inference methods would be required in this case to estimate the time dependent endogenous relationship between the MLTCs [53-55].

Health system costs and out of pocket costs are combined into health care costs in this analysis. A full multi sector analysis would be required to treat these costs as falling on different budgets, consumption in the case of out of pocket costs and the health budget in the case of health system costs [20, 56]. This is potentially an important area of further research.

It should be noted that EVPPI represents the value of eliminating all uncertainty in a group of parameters. Therefore, it represents an upper bound for the value of research. For the task of approximating the relative value of collecting information on different parameters, EVPPI may be a reasonable approximation. However, comparing EVPPI does not consider the (potentially differential) rates at which uncertainty in a given group of parameters are resolved and how this interacts with trial design. An expected value of sample information (EVSI) analysis would be required to fully capture this aspect of research design [15]. This was not carried out here due to computational and evidentiary challenges [57, 58]. As a pragmatic alternative, parameter groups were split into short, medium and long term to capture the impact of trial follow up length, which is a novel approach to addressing the problem.

The analysis was carried out to inform the DiaDeM trial which will be carried out in both Pakistan and Bangladesh. We focused on Pakistan only because of lack of data on patient level and appropriate cost data for routine costs for people with diabetes or costs associated with major cardiovascular events. This is an important limitation of our pre-trial analysis because of potential differences between these countries in for example, costs, health behaviours and care delivery. Following the completion of the DiaDeM, the model will be parameterised for both Bangladesh and Pakistan based on the data collected in the trial.

Though there were more data for Pakistan than Bangladesh, there were still important data limitations for the Pakistani context. Where necessary, data from other countries were used to parameterise the model. For example, the PHQ-9 data were from an Indian study and disease history was imputed based on the relationship between risk factors estimated based on UK data. Data quality was also an issue for sources within Pakistan, for example the cost of care per depressive episode was based on a secondary care facility which may not be representative of practice. For the UKPDS diabetes model the risk of events is based on patient characteristics (e.g. age, smoking status, HbA1c etc). These distribution of characteristics in the population was based on Pakistani data, however, the risk equations linking characteristics to outcomes were based on longitudinal UK data. These assumptions around the generalisability of the evidence to Pakistan are an important limitation of the analysis and highlight the need for more in country research to better inform the model. The model could also be improved in future by further dialogue with local stakeholders and patients. The analysis in this paper does not consider screening or costs involved in identification, therefore the results are implicitly based on the case in which individuals are perfectly identified (all true positives). This also relates to the choice of comparator in the model: usual care. Those who are not identified as having depression will go without any treatment. This no treatment option was not included in the analysis as it was not considered a relevant policy alternative. However, it would be necessary to include in an analysis which included screening which may be relevant for decision makers.

# Conclusion

We found that BA had the potential to be a cost-effective intervention compared to usual care for patients with both depression and diabetes in Pakistan, improving morbidity and mortality and reducing costs. Efforts must be made to keep the BA treatment cost low and uncertainty remains over the impact of depression on diabetes and trajectory of depression. This evidence has helped to inform the design of the DiaDeM intervention by highlighting the value of collecting evidence on the impact of depression on diabetes and the time path of PHQ-9 scores. The model developed will be used to estimate the cost-effectiveness of the intervention following completion of the definitive trial.

# Acknowledgements

Ethics approval was granted by the University of York (HSRGC/2020/409/B), Khyber Medical University (DIR/KMU/UEC/25), Rawalpindi Medical School (242/IREF/RMU/2020) and Diabetic Association from Bangladesh (BADAS-ERC/EC/20/00300) for the study as part of the wider DiaDeM study (publication forthcoming). This study reported in this paper was a secondary analysis of existing data and so informed consent was not required.

Analysis was carried out on the Viking Cluster, which is a high-performance compute facility provided by the University of York. We are grateful for computational support from the University of York High Performance Computing service, Viking and the Research Computing team.

## Funding

This study was carried out under the National Institute of Health Research, Global Health Research project NIHR200806] The University of York is the trial sponsor. The views expressed in this publication are those of the authors and not necessarily those of the NIHR or the UK Department of Health and Social Care. DG received the financial support of the European Union’s Horizon Europe Excellent Science programme under the Marie Skłodowska-Curie Actions Grant Agreement [Grant Agreement No 101081457] and in part from Research Ireland under grant number 13/RC/2073\_P2. The funders had no role in study design, data collection and analysis, decision to publish or preparation of the manuscript.

# Competing interests

RH has recieved honorium for speaking engagements from various pharmaceutical companies. All other authors declare no competing interests relating to this work.

# Author contributor statement

DG, PS and SW carried out the technical analysis. All authors contributed to the methodology and writing of the manuscript. DG is the guarantor.

# References

1. Khunti K, Sathanapally H, Mountain P. Multiple long term conditions, multimorbidity, and co-morbidities: we should reconsider the terminology we use. BMJ: British Medical Journal (Online). 2023;383:p2327.

2. NIHR. NIHR Strategic Framework for Multiple Long-Term Conditions 2020 [Available from: <https://www.nihr.ac.uk/nihr-strategic-framework-multiple-long-term-conditions#:~:text=The%20Multiple%20Long%2Dterm%20Conditions%20(Multimorbidity)%20Strategic%20Framework,-MLTC%2DM%20is&text=The%20NIHR%20will%3A,their%20trajectories%20among%20the%20population>.

3. Pati S, Swain S, Hussain MA, van den Akker M, Metsemakers J, Knottnerus JA, et al. Prevalence and outcomes of multimorbidity in South Asia: a systematic review. BMJ Open. 2015;5(10):e007235.

4. Basit A, Askari S, Zafar J, Riaz M, Fawwad A, Members N. NDSP 06: Prevalence and risk factors for obesity in urban and rural areas of Pakistan: A study from second National Diabetes Survey of Pakistan (NDSP), 2016–2017. Obesity Research & Clinical Practice. 2021;15(1):19-25.

5. Roy T, Lloyd CE. Epidemiology of depression and diabetes: a systematic review. Journal of affective disorders. 2012;142:S8-S21.

6. The Academy of Medical Sciences. Multiple Long-Term Conditions (Multimorbidity): a priority for global health research 2024 [Available from: <https://acmedsci.ac.uk/policy/policy-projects/multimorbidity>.

7. Banstola A, Anokye N, Pokhrel S. The economic burden of multimorbidity: Protocol for a systematic review. Plos one. 2024;19(5):e0301485.

8. ISRCTN registry. ISRCTN40885204. Randomized controlled trial of DiaDeM, an adapted behavioural activation intervention, for people with depression and diabetes in South Asia: BMC; 2023 [Available from: <https://www.isrctn.com/ISRCTN40885204>.

9. Zavala GA, Afaq S, Anas A, Ahmed N, Aslam F, Benkalkar S, et al. Adaptation of a Behavioural Activation Intervention for Depression in People with Diabetes in Bangladesh and Pakistan: DiaDeM Intervention. Global Implementation Research and Applications. 2023;3(1):44-55.

10. Ekers D, Webster L, Van Straten A, Cuijpers P, Richards D, Gilbody S. Behavioural activation for depression; an update of meta-analysis of effectiveness and sub group analysis. PloS one. 2014;9(6):e100100.

11. Uphoff E, Ekers D, Robertson L, Dawson S, Sanger E, South E, et al. Behavioural activation therapy for depression in adults. Cochrane Database of Systematic Reviews. 2020(7).

12. Zavala GA, Jennings HM, Afaq S, Alam A, Ahmed N, Aslam F, et al. Effectiveness and implementation of psychological interventions for depression in people with non-communicable diseases in South Asia: Systematic review and meta-analysis. International Journal of Mental Health. 2023;52(3):260-84.

13. Emmert-Fees KMF, Laxy M, Patel SA, Singh K, Poongothai S, Mohan V, et al. Cost-Effectiveness of a Collaborative Care Model Among Patients With Type 2 Diabetes and Depression in India. Diabetes Care. 2022;46(1):11-9.

14. Kearns B, Rafia R, Leaviss J, Preston L, Brazier JE, Palmer S, et al. The cost-effectiveness of changes to the care pathway used to identify depression and provide treatment amongst people with diabetes in England: a model-based economic evaluation. BMC Health Serv Res. 2017;17(1):78.

15. Briggs A, Sculpher M, Claxton K. Decision modelling for health economic evaluation: Oup Oxford; 2006.

16. Cosh E, Girling A, Lilford R, McAteer H, Young T. Investing in new medical technologies: A decision framework. Journal of Commercial Biotechnology. 2007;13(4):263-71.

17. Girling A, Lilford R, Cole A, Young T. HEADROOM APPROACH TO DEVICE DEVELOPMENT: CURRENT AND FUTURE DIRECTIONS. Int J Technol Assess Health Care. 2015;31(5):331-8.

18. Rothery C, Strong M, Koffijberg HE, Basu A, Ghabri S, Knies S, et al. Value of Information Analytical Methods: Report 2 of the ISPOR Value of Information Analysis Emerging Good Practices Task Force. Value Health. 2020;23(3):277-86.

19. Fenwick E, Steuten L, Knies S, Ghabri S, Basu A, Murray JF, et al. Value of Information Analysis for Research Decisions-An Introduction: Report 1 of the ISPOR Value of Information Analysis Emerging Good Practices Task Force. Value Health. 2020;23(2):139-50.

20. Walker S, Griffin S, Asaria M, Tsuchiya A, Sculpher M. Striving for a societal perspective: a framework for economic evaluations when costs and effects fall on multiple sectors and decision makers. Applied health economics and health policy. 2019;17(5):577-90.

21. Wilkinson T, Sculpher MJ, Claxton K, Revill P, Briggs A, Cairns JA, et al. The International Decision Support Initiative Reference Case for Economic Evaluation: An Aid to Thought. Value Health. 2016;19(8):921-8.

22. Clarke PM, Gray AM, Briggs A, Farmer AJ, Fenn P, Stevens RJ, et al. A model to estimate the lifetime health outcomes of patients with type 2 diabetes: the United Kingdom Prospective Diabetes Study (UKPDS) Outcomes Model (UKPDS no. 68). Diabetologia. 2004;47(10):1747-59.

23. Hayes AJ, Leal J, Gray AM, Holman RR, Clarke PM. UKPDS outcomes model 2: a new version of a model to simulate lifetime health outcomes of patients with type 2 diabetes mellitus using data from the 30 year United Kingdom Prospective Diabetes Study: UKPDS 82. Diabetologia. 2013;56(9):1925-33.

24. Leal J, Alva M, Gregory V, Hayes A, Mihaylova B, Gray AM, et al. Estimating risk factor progression equations for the UKPDS Outcomes Model 2 (UKPDS 90). Diabetic Medicine. 2021;38(10):e14656.

25. Siddiqui F, Aslam F, Ahmed N, Afaq S, Khan A, Keding A, et al. Effectiveness and cost-effectiveness of a culturally tailored Behavioural Activation intervention (DiaDeM) for treating Depression in Type 2 Diabetes: Protocol for a parallel arm, multi-country, randomised controlled trial in South Asia. medRxiv. 2023:2023.08.11.23294009.

26. R Core Team. R: A language and environment for statistical computing. R Foundation for Statistical Computing. Vienna, Austria: R Foundation for Statistical Computing; 2021.

27. Gupta, Willis M, Johansen P, Nilsson A, Shah M, Mane A, et al. Long-Term Clinical Benefits of Canagliflozin 100 mg Versus Sulfonylurea in Patients With Type 2 Diabetes Mellitus Inadequately Controlled With Metformin in India. Value Health Reg Issues. 2019;18:65-73.

28. Tosh J, Kearns B, Brennan A, Parry G, Ricketts T, Saxon D, et al. Innovation in health economic modelling of service improvements for longer-term depression: demonstration in a local health community. BMC health services research. 2013;13(1):1-11.

29. Ali MK, Chwastiak L, Poongothai S, Emmert-Fees KMF, Patel SA, Anjana RM, et al. Effect of a Collaborative Care Model on Depressive Symptoms and Glycated Hemoglobin, Blood Pressure, and Serum Cholesterol Among Patients With Depression and Diabetes in India: The INDEPENDENT Randomized Clinical Trial. JAMA. 2020;324(7):651-62.

30. Schmitt A, Bendig E, Baumeister H, Hermanns N, Kulzer B. Associations of depression and diabetes distress with self-management behavior and glycemic control. Health Psychol. 2021;40(2):113-24.

31. van Steenbergen-Weijenburg KM, van Puffelen AL, Horn EK, Nuyen J, van Dam PS, van Benthem TB, et al. More co-morbid depression in patients with Type 2 diabetes with multiple complications. An observational study at a specialized outpatient clinic. Diabet Med. 2011;28(1):86-9.

32. Cuijpers P, Quero S, Noma H, Ciharova M, Miguel C, Karyotaki E, et al. Psychotherapies for depression: a network meta‐analysis covering efficacy, acceptability and long‐term outcomes of all main treatment types. World Psychiatry. 2021;20(2):283-93.

33. Kennedy MC, O'Hagan A. Bayesian calibration of computer models. Journal of the Royal Statistical Society: Series B (Statistical Methodology). 2001;63(3):425-64.

34. Alarid-Escudero F, MacLehose RF, Peralta Y, Kuntz KM, Enns EA. Nonidentifiability in model calibration and implications for medical decision making. Medical Decision Making. 2018;38(7):810-21.

35. Baqai Institute of Diabetology & Endocrinology. Patient registry 2023 [Available from: <https://bide.edu.pk/>.

36. Bourne R, Dineen B, Jadoon Z, Lee PS, Khan A, Johnson GJ, et al. Outcomes of cataract surgery in Pakistan: results from the Pakistan National Blindness and Visual Impairment Survey. British journal of ophthalmology. 2007;91(4):420-6.

37. Gupta, Singh K, Shivashankar R, Singh K, Vamadevan A, Mohan V, et al. Healthcare utilisation and expenditure patterns for cardio-metabolic diseases in South Asian cities: the CARRS Study. BMJ open. 2020;10(9):e036317.

38. Alva ML, Gray A, Mihaylova B, Leal J, Holman RR. The impact of diabetes-related complications on healthcare costs: new results from the UKPDS (UKPDS 84). Diabet Med. 2015;32(4):459-66.

39. Malik MA, Khan MM. Economic Burden of Mental Illnesses in Pakistan. J Ment Health Policy Econ. 2016;19(3):155-66.

40. Salomon JA, Haagsma JA, Davis A, de Noordhout CM, Polinder S, Havelaar AH, et al. Disability weights for the Global Burden of Disease 2013 study. The Lancet Global Health. 2015;3(11):e712-e23.

41. Organisation WH. Global burden of disease 2019 [Available from: <http://ghdx.healthdata.org/gbd-results-tool>.

42. Lomas J, Claxton K, Ochalek J. Accounting for country-and time-specific values in the economic evaluation of health-related projects relevant to low-and middle-income countries. Health policy and planning. 2022;37(1):45-54.

43. Ochalek J, Lomas J. Reflecting the Health Opportunity Costs of Funding Decisions Within Value Frameworks: Initial Estimates and the Need for Further Research. Clin Ther. 2020;42(1):44-59.e2.

44. Ochalek J, Lomas J, Claxton K. Estimating health opportunity costs in low-income and middle-income countries: a novel approach and evidence from cross-country data. BMJ Global Health. 2018;3(6):e000964.

45. World Bank. Population, total for Pakistan 2020 [Available from: <https://data.worldbank.org/country/pakistan>.

46. Uphoff, Newbould L, Walker I, Ashraf N, Chaturvedi S, Kandasamy A, et al. A systematic review and meta-analysis of the prevalence of common mental disorders in people with non-communicable diseases in Bangladesh, India, and Pakistan. J Glob Health. 2019;9(2):020417.

47. Akhtar S, Nasir JA, Abbas T, Sarwar A. Diabetes in Pakistan: A systematic review and meta-analysis. Pak J Med Sci. 2019;35(4):1173-8.

48. Patel V, Weobong B, Weiss HA, Anand A, Bhat B, Katti B, et al. The Healthy Activity Program (HAP), a lay counsellor-delivered brief psychological treatment for severe depression, in primary care in India: a randomised controlled trial. The Lancet. 2017;389(10065):176-85.

49. Culyer AJ. Cost-effectiveness thresholds in health care: a bookshelf guide to their meaning and use. Health Economics, Policy and Law. 2016;11(4):415-32.

50. Kuyken W, Byford S, Taylor RS, Watkins E, Holden E, White K, et al. Mindfulness-based cognitive therapy to prevent relapse in recurrent depression. Journal of consulting and clinical psychology. 2008;76(6):966.

51. Simon GE, Von Korff M, Ludman EJ, Katon WJ, Rutter C, Unützer J, et al. Cost-effectiveness of a program to prevent depression relapse in primary care. Medical Care. 2002;40(10):941-50.

52. Rodgers M, Asaria M, Walker S, McMillan D, Lucock M, Harden M, et al. The clinical effectiveness and cost-effectiveness of low-intensity psychological interventions for the secondary prevention of relapse after depression: a systematic review. Health Technol Assess. 2012;16(28):1-130.

53. Hernán, Robins. Causal Inference: What If.: Boca Raton: Chapman & Hall/CRC; 2020.

54. Gran JM, Røysland K, Wolbers M, Didelez V, Sterne JA, Ledergerber B, et al. A sequential Cox approach for estimating the causal effect of treatment in the presence of time‐dependent confounding applied to data from the Swiss HIV Cohort Study. Statistics in medicine. 2010;29(26):2757-68.

55. Naimi AI, Cole SR, Kennedy EH. An introduction to g methods. International journal of epidemiology. 2017;46(2):756-62.

56. Ramponi F, Walker S, Griffin S, Parrott S, Drummond C, Deluca P, et al. Cost-effectiveness analysis of public health interventions with impacts on health and criminal justice: An applied cross-sectoral analysis of an alcohol misuse intervention. Health Econ. 2021;30(5):972-88.

57. Heath A, Strong M, Glynn D, Kunst N, Welton NJ, Goldhaber-Fiebert JD. Simulating Study Data to Support Expected Value of Sample Information Calculations: A Tutorial. Medical Decision Making. 2022;42(2):143-55.

58. Strong M, Oakley JE, Brennan A, Breeze P. Estimating the Expected Value of Sample Information Using the Probabilistic Sensitivity Analysis Sample: A Fast, Nonparametric Regression-Based Method. Med Decis Making. 2015;35(5):570-83.

# Figure legends

FIG1

Counterfactual PHQ-9 path for a single individual with usual care (black line) or usual care plus BA (blue line). In both cases the individual begins the model at the start of a depressive episode and has two additional episodes at month 15 and month 55. Between episodes the individual recovers gradually, with the rate of recovery being twice as fast with BA. The PHQ-9 score at the beginning of a new episode is determined by the baseline PHQ-9 score and the PHQ-9 score before the episode began. PHQ-9 score cannot go above 27 or below 0. PHQ-9 = Patient Health Questionnaire-9; BA = Behavioural activation.

FIG2

Two-way interaction between diabetes and depression. HbA1c = hemoglobin A1c, PHQ-9 = Patient Health Questionnaire-9.