Rationale and protocol for estimating the economic value of a multicomponent quality improvement strategy for diabetes care in South Asia.

Kavita Singh, Public Health Foundation of India
Mohammed Ali, Emory University
Raji Devarajan, Public Health Foundation of India
Roopa Shivashankar, Centre for Chronic Disease Control
Dimple Kondal, Public Health Foundation of India
Vamadevan S. Ajay, Public Health Foundation of India
V. Usha Menon, Amrita Institute of Medical Sciences
Premlata K. Varthakavi, BYL Nair Charity Hospital
Vijay Viswanathan, MV Hospital for Diabetes & Diabetes Research Centre
Mala Dharmalingam, Bangalore Endocrinology & Diabetes Research Centre

Only first 10 authors above; see publication for full author list.

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Rationale and protocol for estimating the economic value of a multicomponent quality improvement strategy for diabetes care in South Asia

Kavita Singh1,16*, Mohammed K. Ali2, Raji Devarajan1,16, Roopa Shivashankar16, Dimple Kondal1,16, Vamadevan S. Ajay1,16, V. Usha Menon3, Premala K. Varthakavi3, Vijay Viswanathan5, Mala Dharmalingam6, Ganapati Bantwal7, Rakesh Kumar Sahay8, Muhammad Qamar Masood9, Rajesh Khadgawat10, Ankush Desai11, Dorairaj Prabhakaran12,16,17, K. M. Venkat Narayan13, Victoria L. Phillips14, Nikhil Tandon15 and On behalf of the CARRS Trial Group

Abstract

Background: Economic dimensions of implementing quality improvement for diabetes care are understudied worldwide. We describe the economic evaluation protocol within a randomised controlled trial that tested a multi-component quality improvement (QI) strategy for individuals with poorly-controlled type 2 diabetes in South Asia.

Methods/design: This economic evaluation of the Centre for Cardiometabolic Risk Reduction in South Asia (CARRS) randomised trial involved 1146 people with poorly-controlled type 2 diabetes receiving care at 10 diverse diabetes clinics across India and Pakistan. The economic evaluation comprises both a within-trial cost-effectiveness analysis (mean 2.5 years follow up) and a microsimulation model-based cost-utility analysis (life-time horizon). Effectiveness measures include multiple risk factor control (achieving HbA1c < 7% and blood pressure < 130/80 mmHg and/or LDL-cholesterol < 100 mg/dl), and patient reported outcomes including quality adjusted life years (QALYs) measured by EQ-5D-3 L, hospitalizations, and diabetes related complications at the trial end. Cost measures include direct medical and non-medical costs relevant to outpatient care (consultation fee, medicines, laboratory tests, supplies, food, and escort/accompanying person costs, transport) and inpatient care (hospitalization, transport, and accompanying person costs) of the intervention compared to usual diabetes care. Patient, healthcare system, and societal perspectives will be applied for costing. Both cost and health effects will be discounted at 3% per year for within trial cost-effectiveness analysis over 2.5 years and decision modelling analysis over a lifetime horizon. Outcomes will be reported as the incremental cost-effectiveness ratios (ICER) to achieve multiple risk factor control, avoid diabetes-related complications, or QALYs gained against varying levels of willingness to pay threshold values. Sensitivity analyses will be performed to assess uncertainties around ICER estimates by varying costs (95% CIs) across public vs. private settings and using conservative estimates of effect size (95% CIs) for multiple risk factor control. Costs will be reported in US$ 2018.

(Continued on next page)
Background
Diabetes is one of the fastest growing public health problems with huge financial burdens. The global costs of diabetes were US$ 1.31 trillion (1.8% of global GDP) in 2015 [1]. A 2018 systematic review found that annual costs of diabetes care (out of pocket medical expenditure) in South Asia ranged between US$ 575 to US $1216 per person [2]. Diabetes is a progressive disease which requires increasingly more clinic visits, laboratory tests, and patients need to engage with the healthcare system and providers over years for better management of diabetes which can arrest disease progression. However, current chronic care for diabetes is sub-optimal, costly, and lower socioeconomic status or uninsured individuals may be more likely to experience poor control [3–7].

Several barriers at the patient- (e.g., low motivation, financial barriers), provider- (e.g., inertia to intensify treatments), and system-level (e.g., complicated and/or fragmented care system), individually or together, cause patient and system “fatigue” and disrupt achievement of diabetes care goals [8–10]. In the Centre for Cardiovascular Risk Reduction in South Asia (CARRS) Trial, we targeted different levels of barriers together (e.g., patient motivation and provider inertia) [9, 11] and demonstrated sustainable and larger improvements in outcomes and satisfaction for people with diabetes with a multicomponent strategy of decision support-electronic health records (DS-EHR) and non-physician care coordinators (CC), compared to usual diabetes care [12].

However, enhancements or changes to the status quo of care delivery come at a cost, and in order to formulate useful recommendations for practicing clinicians, health systems, payers (health insurance, governments, patients paying out-of-pocket), and policymakers, there is an imperative to assess the value of investing in quality improvement (QI) care models. Knowing the upfront costs is also necessary to guide decision makers as they consider implementation of QI interventions in clinical care.

A 2018 systematic review of economic evaluations of QI interventions for glycaemic control among adults with type 1 or type 2 diabetes from high income countries found that multifaceted QI interventions that lower HbA1c was good value for money versus usual care, depending on society’s willingness to pay [13]. However, in our review of cost-effectiveness of interventions to control cardiovascular diseases and diabetes mellitus, we found a scarcity of cost-effectiveness studies related to QI interventions for diabetes care in South Asia [14]. Here, we describe the economic evaluation protocol to assess the within-trial cost-effectiveness and broader societal value of the CARRS diabetes care model consisting of DS-EHR and non-physician CCs compared to usual diabetes care.

Methods/Design of Economic Evaluation
Overview
The objectives of the economic evaluation are to assess: a) the incremental cost of delivering multicomponent QI interventions compared to usual diabetes care in tertiary care settings over a period of 2.5 years; b) whether the intervention provides value for money (cost-effectiveness) to patients, healthcare systems and society than usual care, and if so; c) the extent of uncertainty over the cost-effectiveness of the intervention and value of conducting further research to reduce this uncertainty.

The CARRS Trial’s economic evaluation will follow standard international methodological guidelines [15–18]. Given, more than 80% of medical expenses in India and Pakistan are out-of-pocket expenditures borne by the patient, we will apply the patient viewpoint as the predominant perspective, in addition to healthcare system and societal perspectives for costing resource use. Cost data will be reported in 2018 United States Dollars (US$). Both cost and health effects will be discounted at 3% per year as per the World Health Organization’s (WHO) guidelines for conducting economic evaluations in developing countries.

The CARRS trial and study population
The CARRS Trial randomised 1146 eligible patients with poorly controlled type 2 diabetes (HbA1c > 8% and SBP > 140 mmHg or LDLc> 130 mg/dl) to intervention (n = 575) or usual care (n = 571) across 10 diverse diabetes clinics in India and Pakistan. At baseline, participants’ mean age was 54 years, 45% were males, mean HbA1c was 9.9%, LDLc 123.2 mg/dl, BP 144.2/82.3 mmHg, and median duration of diabetes was 7 years [12].
**Intervention and comparator**

Detailed information about the CARRS-Trial intervention and protocol has been published previously [19]. Briefly, the CARRS intervention consisted of DS-EHRs to enhance physicians’ responsiveness to consider treatment modification and non-physician CCs to support patients in their adherence to prescribed therapies. The DS-EHR stored all consultation, laboratory, self-care, and diabetes related complications data for patients in one easily accessible web portal to monitor patient progress; and provided decision-support system (DSS) prompts to facilitate achievement of guideline-recommended glycemic, blood pressure, and lipid goals. The CCs fully managed the DS-EHR data-entry for intervention group participants and all communication of DSS prompts to the physician during consultations via print-out or electronic display. Physicians could, at their discretion, accept or reject DSS prompts and modify treatment plans based on clinical judgment, so long as justification was provided.

The intervention was compared with usual diabetes care at nine clinics/hospitals across India and one site in Pakistan. Figure 1 demonstrates the study flow.

**Effectiveness measures**

To evaluate incremental effectiveness, we will compare the proportions of intervention and control arm participants achieving multiple risk factor control defined as HbA1c < 7% and BP < 130/80 mmHg or LDLc< 100 mg/dl (and < 70 mg/dl for those with history of cardiovascular disease). Data on health-related quality of life (EQ5D-3 L); new-onset cardiovascular events, new onset microvascular events, and other hospitalizations would also be used.

The CARRS Trial is currently ongoing and we will project cardiovascular and microvascular outcomes using proxy indicators (intermediate risk factors: HbA1c, BP, LDLc). Relative risk reductions for major adverse cardiovascular events with intervention or comparator will also be calculated separately for each participant, using United Kingdom Prospective Diabetes Study (UKPDS) Outcomes Models 2 which has been validated for use in South Asians [20]. Table 1 summarizes the study outcomes (effectiveness measures) to be considered in the cost-effectiveness analysis.

**Resource use and cost data**

Resource utilization and costs will be estimated using data from the CARRS Trial population (1146 participants). The study paid the costs of annual laboratory investigations, but patients had to bear the cost of clinic visits and laboratory tests for regular follow-up visits or any other interim clinic visits, tests, medication changes, or procedures advised by the treating physician. CARRS Trial data will be extracted from clinic and study records for the following: medication use, laboratory tests, consultations with healthcare professionals (outpatient attendance for diabetes); preventive screening (eye examination, foot examination, ECG, microalbuminuria test), emergency department attendances (when not admitted to hospital); and serious adverse events (including all hospital admissions).

Patients’ self-reported expenditures and costs of outpatient visits and hospitalizations related to diabetes complications will be extracted from the trial annual visit case report forms (CRF). Out-of-pocket expenses reported by the patients will permit estimation of economic value from the patient’s perspective.

To estimate value from a healthcare system perspective, unit costs for outpatient visits and in-patient hospitalizations, and processes of care measures including preventive examinations will be obtained from participating hospitals. For treatment of cardiovascular and microvascular events, we will extract detailed information concerning diagnosis; length of hospital stay; diagnostic/therapeutic procedures and any ongoing treatment and support. Additionally, the unit price of medications will be obtained from the PharmaTrac database for January 2014 [21]. PharmaTrac provides the market retail price (MRP) of all drugs by drug class, brand name, generic composition, formulation (oral/injectables), dose, and packs being sold in India. PharmaTrac has an extensive coverage of drug retailers and is believed to be a reliable source to estimate unit cost of drug prices in India. The IMS Health drug database will be used to estimate drug prices in Pakistan.

To estimate costs from the societal perspective, indirect costs due to lost productivity (number of work days missed due to out-patient or in-patient care) will be valued using the human capital approach [22]. Finally, total costs over the trial period and annual cost per patient (both undiscounted and discounted) will be estimated for individual patients by multiplying resource use by unit costs.

**Intervention costs**

Intervention development and delivery costs will be derived from the CARRS Trial expense records (accounts register) and will be estimated from the health system perspective. Intervention costs include DS-EHR development, implementation, and maintenance costs; intervention training; care coordinator salary; and the incremental health care costs associated with the intervention delivery (i.e. the costs of additional medicines, additional clinic visits that patients bear and whether it is different between the treatment groups). These costs will be calculated as average costs of implementation per person and exclude any research specific costs. The cost estimates assume that the DS-EHR is implemented in a relatively large tertiary care hospital having additional resource facilities to implement the intervention (i.e. workspace for the care coordinator, and access to internet service providers is considered a maintenance
cost). DS-EHR development and maintenance costs will include software programmer’s time, expert consultant/physicians time in developing and reviewing the diabetes management algorithm. DS-EHR implementation cost will include care coordinators and site physician’s time in entering patient details in the EHR system and review of software generated diabetes management plans, respectively. Intervention training costs include training materials, the time of the trainers and the staff participating in the training, and training for physicians to use the DS-EHR algorithms. These costs will be estimated using the study’s accounting data. Training material and time costs will be estimated from the first year of the intervention. Tables 2 and 3 present an overview of cost measures, health service use, and source of data.

**Within trial cost-effectiveness analysis**

Based on estimates of between-group differences in mean healthcare costs and outcomes (adjusting for differences in baseline characteristics) over the study period, we will estimate the following incremental cost-effectiveness ratios (ICERs):

- Incremental cost per primary outcome achieved (i.e. multiple risk factor control: HbA1c < 7% and BP < 130/80 mmHg and/or LDLc < 100 mg/dl)
than 10% of a domain/variable [29] missing cost values if missing data accounts for more we will use multiple imputation approaches to replace address potential biases due to incomplete follow-up, imputing values within each dimension [25] ing data; that is, missing data will be handled by

time, we will follow the developer
tion, we will follow international guidelines for verifica-

d and validation of decision models [32].

### Table 1 Overview of the effectiveness measures

<table>
<thead>
<tr>
<th>Measure</th>
<th>Means of collection</th>
<th>Timing of collection</th>
<th>Source of data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multiple risk factor control (HbA1c &lt; 7% and BP &lt; 130/80 mmHg or LDLc &lt; 100 mg/dl)</td>
<td>Blood test + BP measurement using digital BP monitor</td>
<td>Baseline: Prior to intervention deliveryFollow-up: Annual visits post intervention delivery</td>
<td>Trial eCRF (Form C, E, F)</td>
</tr>
<tr>
<td>Single risk factor control</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HbA1c (1% point reduction)</td>
<td>Fasting blood test</td>
<td>Baseline: Prior to intervention deliveryFollow-up: Annual visits post intervention delivery</td>
<td>Trial eCRF (Form C, E, F)</td>
</tr>
<tr>
<td>SBP (5 mmHg reduction)</td>
<td>BP measurement using digital BP monitor (Ommron-T9P)</td>
<td>Baseline: Prior to intervention deliveryFollow-up: Annual visits post intervention delivery</td>
<td>Trial eCRF (Form C, E, F)</td>
</tr>
<tr>
<td>DBP (5 mmHg reduction)</td>
<td>BP measurement using digital BP monitor (Ommron-T9P)</td>
<td>Baseline: Prior to intervention deliveryFollow-up: Annual visits post intervention delivery</td>
<td>Trial eCRF (Form C, E, F)</td>
</tr>
<tr>
<td>LDLc (10 mg/dl reduction)</td>
<td>Fasting blood test</td>
<td>Baseline: Prior to intervention deliveryFollow-up: Annual visits post intervention delivery</td>
<td>Trial eCRF (Form C, E, F)</td>
</tr>
<tr>
<td>Major adverse cardiovascular events</td>
<td>Self-reported by patient and physician verified</td>
<td>Follow-up: All study related and non-study related clinic visits</td>
<td>Trial eCRF (form X)</td>
</tr>
<tr>
<td>Diabetes related micro-vascular complications</td>
<td>Self-reported by patient and physician verified</td>
<td>Follow-up: All study related and non-study related clinic visits</td>
<td>Trial eCRF (form X)</td>
</tr>
<tr>
<td>Quality adjusted life years</td>
<td>EQ5D-3 L</td>
<td>Baseline: Prior to intervention deliveryFollow-up: Annual visits post intervention delivery</td>
<td>Trial eCRF (Form C, E, F)</td>
</tr>
</tbody>
</table>

HbA1c: Glycated haemoglobin, SBP: Systolic blood pressure, LDLc: Low-density lipoprotein cholesterol, eCRF: Electronic case report form, EQSD-3 L: European Quality of Life five dimension 3 levels, BP Blood pressure, mg/dl: Milligrams per deciliter, mmHg: Millimeter of mercury

- Incremental cost per unit reduction in single risk factors: HbA1c (1% point reduction), SBP (5 mmHg reduction), and LDLc (10 mg/dl reduction)
- Incremental cost per quality adjusted life years (QALYs) gained

Non-parametric bootstrapping will be used to report 95% confidence intervals around the ICER estimates [23]. ICERs will be reported in US$ 2018. Cost effectiveness acceptability curves against a wide range of willingness to pay values will be presented [24]. Cost-effectiveness results will be also presented by major sub-groups: age, gender, education, income level, types of health setting (public, private or semi-private) and history of macro- and micro-vascular complications.

**Decision-modeling based cost-utility analysis**

A decision-analytic microsimulation model will be developed to evaluate long-term costs and health consequences of delivering care for people with type 2 diabetes using a multicomponent QI strategy rather than current standard care approaches. A microsimulation model is chosen as it is very flexible and can reflect complex treatment pathways and relationships between individuals’ characteristics, histories, and outcomes; it can be used to examine the impact of real resource constraints within a healthcare system.

The microsimulation decision model will be implemented using appropriate software: STATA or a programming language (e.g. R). To assure the credibility of our model, we will follow international guidelines for verification and validation of decision models [32].

**Missing data**

The CARRS Trial has a minimal loss to follow-up including consent withdrawals and deaths at 2.5 years (9.2%) but, if needed; multiple imputation approaches will be used to handle missing outcomes data [12]. For EQ5D-3 L scores, which will be used for QALY estimation, we will follow the developer’s guideline for missing data; that is, missing data will be handled by imputing values within each dimension [25–28]. To address potential biases due to incomplete follow-up, we will use multiple imputation approaches to replace missing cost values if missing data accounts for more than 10% of a domain/variable [29–31]. Since cost data are unlikely to be normally distributed, [29] we will use the multiple imputation chained equations approach to impute missing cost data. Costs will be imputed at the total cost level [29].

**Model analysis**

All analyses will compare results for the CARRS multicomponent QI care delivery model versus usual diabetes care. In the CARRS Trial microsimulation model, costs and QALYs will be recorded for each individual and an average cost and QALY for the simulated population will be estimated. The microsimulation model will be run twice, once to simulate costs and QALYs under usual care and the other to simulate costs and QALYs under the intervention scenario (multicomponent QI strategy). Individuals representing the CARRS trial inclusion criteria will enter the model and their baseline risk for CVD events and diabetes-related microvascular complications will be estimated using the UKPDS Outcomes Model 2 algorithm. Costs and QALYs will be recorded for each event (including adverse events). Individuals can experience more than one event (model run for lifetime horizon) and patient
<table>
<thead>
<tr>
<th>Type of cost</th>
<th>Level</th>
<th>Expense type</th>
<th>Cost component</th>
<th>Means of collection</th>
<th>Timing of collection</th>
<th>Source of data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct</td>
<td>Intervention</td>
<td>Fixed cost</td>
<td>Software development</td>
<td>Trial records</td>
<td>After completion of software development</td>
<td>Developers of software: DS-EHR</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fixed cost</td>
<td>Training of physicians and care coordinators</td>
<td>Trial records</td>
<td>After completion of training</td>
<td>Trial Team</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fixed cost</td>
<td>Laptop</td>
<td>Trial records</td>
<td>Baseline (at the start of the trial)</td>
<td>Trial records</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fixed cost</td>
<td>Mobile phone</td>
<td>Trial records</td>
<td>Baseline (at the start of the trial)</td>
<td>Trial records</td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Care coordinator’s salary</td>
<td>Trial records</td>
<td>Monthly</td>
<td>Payment invoice</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Three-monthly laboratory tests</td>
<td>Interview with patients</td>
<td>Annual</td>
<td>Self-reported invoice</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Internet</td>
<td>Trial records</td>
<td>Annual</td>
<td>Payment invoice</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Communication cost</td>
<td>Trial records</td>
<td>Annual</td>
<td>Payment invoice</td>
<td></td>
</tr>
<tr>
<td>Fixed cost</td>
<td>Clinic/Hospital</td>
<td>Software maintenance</td>
<td>Trial records</td>
<td>Annual</td>
<td>Payment invoice</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Physician’s time</td>
<td>Interview with patients and physicians</td>
<td>Annual</td>
<td>Self-reported by physicians</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Resource use for patient management: telephone calls, letters, team meetings</td>
<td>Interview with physicians and hospital administrators</td>
<td>Annual</td>
<td>Self-reported by administrators</td>
<td></td>
</tr>
<tr>
<td>Patient</td>
<td>Variable cost</td>
<td>Medications</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Medical supplies (glucose strips, gauze, sterile solution, etc.)</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Laboratory tests</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Diagnostics</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Preventative screening (ECG, eye exam, foot exam, dental exam, etc.)</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Outpatient visits (consultation fee)</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Transportation</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Food (personal)</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Additional cost for escort</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Other out of pocket expenses</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>In-patient hospitalization</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Procedures</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Indirect</td>
<td>Patient</td>
<td>Variable cost</td>
<td>Lost-productivity</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Work days lost due to outpatient visit</td>
<td>Interview with patients + Trial eCRF</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td></td>
<td>Work days lost due to</td>
<td>Interview with patients</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
</tbody>
</table>
characteristics such as age and history of previous events, such as a stroke or diabetic retinopathy, will be updated as the model is being run, with ensuing reflective increases in the risk of an event. The simulation model will run for a sufficient number of iterations to provide stable results. If there is a trade-off between costs and health effects (higher costs and better health outcomes for the CARRS intervention, or vice versa), the incremental cost per cardiovascular event averted, incremental cost per diabetes-related microvascular complication averted, and incremental cost per quality adjusted life year (QALY) gained will be reported.

### Table 2 Overview of cost measures (Continued)

<table>
<thead>
<tr>
<th>Type of cost</th>
<th>Level</th>
<th>Expense type</th>
<th>Cost component</th>
<th>Means of collection</th>
<th>Timing of collection</th>
<th>Source of data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variable cost</td>
<td>loss of concentration</td>
<td>+ Trial eCRF</td>
<td>Interview with patients</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
<tr>
<td>Variable cost</td>
<td>loss of function (health utility index)</td>
<td>+ Trial eCRF</td>
<td>Interview with patients</td>
<td>Annual</td>
<td>Self-reported by patient in eCRF</td>
<td></td>
</tr>
</tbody>
</table>

DS-EHR Decision-support Electronic Health Records, ECG Electrocardiogram, eCRF Electronic case report form

Projections of cost-effectiveness estimates over a lifetime horizon will be made for India and Pakistan.

### Sensitivity analysis

Several one-way sensitivity analyses will be carried out to estimate the uncertainties around ICERs. First, to address the uncertainty around the ICER relating to external validity, we will carry out sensitivity analyses on the most important cost drivers (medications, hospitalizations, and consultation fees) to assess the impact of protocol-driven healthcare use. Second, total cost will be calculated with and without the costs of developing the intervention (DS-EHR) to ascertain whether an increased cost in the intervention arm could be explained by costs for some of the components of the intervention. Lastly, sensitivity analyses would vary the effectiveness of the intervention in trial vs. non-trial settings based on the lower and upper limit of 95% confidence intervals (CI) of the effect estimates. Results of probabilistic sensitivity analyses will be presented using a scatter plot of points on the cost-effectiveness plane – illustrating the possible ranges of estimates of incremental costs and incremental QALYs [24].

### Discussion

The publication and peer-review of economic evaluation protocols alongside clinical trials is recommended to increase transparency and minimise bias [33]. Here, we describe the protocol of an economic evaluation of a multi-component QI strategy compared to usual diabetes care in South Asia from patient, healthcare system, and societal perspectives. There are very few economic evaluations of QI strategies for chronic disease management in South Asia [34] or in LMICs in general, and so this report fills a gap. Following internationally recognised guidelines [15], this protocol serves to heighten the transparency of our economic evaluation approach.

Economic evaluations from high-income countries demonstrate that multifactorial QI strategies are cost-effective. For example, the STENO-2 study showed that, from a health care payer perspective in Denmark, intensive multifactorial intervention was more cost-effective than conventional treatment (ICER: €2538 or US$ 2954 per QALY gained) over a lifetime horizon [35]. Increased costs with intensive treatment were due to increased pharmacy and consultation costs. However, this also resulted in more
QALYs gained for intensive treatment versus conventional treatment (+1.66 QALYs). The ADDITION-UK trial based cost-effectiveness analysis comparing intensive versus conventional treatment demonstrated an ICER of £71,232 (US$93,566)/QALY, £28,444 (US$37,362)/QALY, and £27,549 (US$36,186)/QALY over 10-, 20-, and 30-year time horizons respectively [36]. Given the United Kingdom's willingness-to-pay thresholds in patients with diabetes, intensive treatment was of borderline cost-effectiveness over a time horizon of ≥20 years. The estimates of cost-effectiveness from the CARRS Trial will provide much needed data on whether a simple multifactorial intervention can improve health outcomes with modest increases in costs in resource-constrained settings.

To enhance external validity, it is recommended that evaluations using randomised controlled trials should identify threats to validity such as recruitment/selection bias, protocol-driven utilisation, and enhanced compliance [34, 37]. Regarding recruitment biases, the CARRS Trial’s multicentre approach and inclusion of public, private, and semi-private practices increases the generalisability and transferability of our economic evaluation findings [38]. Further, we will extrapolate the decision analytic microsimulation model beyond the within-trial analysis by using a sample population of poorly controlled type 2 diabetes patients in India / Pakistan stratified by age-group, gender, and location. Also, although Markov models can also be adapted for this purpose, microsimulation models are better suited for analysis of a mixed population with both incident and prevalent diabetes complications (cardiovascular diseases and microvascular events) [39].

This study has several strengths. First, the economic evaluation protocol follows recognised international guidelines to design and report on the relative costs and benefits of an intervention tested in a randomised trial [15, 37]. Second, the economic evaluation will include individual patient-level data over a lengthy 2.5 years of follow up, which are preferable for economic evaluations [15]. Importantly, these patient-level data include objective measures of health outcomes, health service use, and medicine use, all obtained during the trial [40]. Reliable economic evaluations are crucial to shape healthcare policy, in particular when the possibility of bias in economic evidence has been minimised by randomisation [40]. Third, our cost-effectiveness results will also provide a range of values for both the cost of achieving multiple risk factor targets but also costs to achieve single and combined risk factor improvements from poorly controlled baseline values (mean baseline HbA1c = 9.9) from various perspectives (patient, healthcare system and societal). Given a large proportion of healthcare in South Asia is paid for out-of-pocket, our economic analyses consider that scenario explicitly with a patient perspective analysis. Fourth, our proposed micro-simulation model based on UKPDS Outcomes Model 2 will enable long-term cost-effectiveness analysis and a population budget impact analysis which will provide cross-sectional estimates of population impact by year for planning purposes and scalability of the intervention.

This study has a few noteworthy limitations. First, reliance on patient self-reported out-of-pocket medical costs may impact the validity of study results. A 2016 systematic review of validated self-reported questionnaires to measure resource utilisation and costs in economic evaluation concluded that self-reported questionnaires had good agreement with administrative data and are a valid method of collecting data on health resource utilisation and associated costs [41]. However, to overcome any reporting bias in self-reported costs data, a sub-set of self-reported costs will be verified against the administrative data and we will carry out several one-way and probabilistic sensitivity analyses around the self-reported costs in the microsimulation model to estimate the confidence in the reported ICER values. Another limitation of the proposed evaluation is that in India and Pakistan, there is not an explicit willingness-to-pay threshold for reduction of cardiovascular risk in people with diabetes, or an explicit willingness-to-pay threshold for cost per unit reductions in CVD risk factors. As such, it is hard to declare how patients value the intervention. The Commission for Macroeconomics and Health recommends using a threshold of 1-3x GDP per capita per QALY gained or DALYs averted to define cost-effectiveness of a new intervention when conducting global or regional economic evaluations [16, 42]. Although arbitrary, we will use this threshold as it has been used previously and has some philosophical underpinnings [43]. We will apply these and then perform a sensitivity analysis for the main economic outcome and present the cost-effectiveness results on a cost-effectiveness acceptability curve considering a wide range of willingness to pay values. Common to all cost-effectiveness analyses conducted alongside randomised trials, external validity of the results may be influenced by restrictive inclusion criteria and protocol-driven resource use, among other factors [15, 37]. Therefore, we will conduct a range of sensitivity analyses around key variables (cost drivers, total cost calculated with and without the cost of the intervention development, patient characteristics, and effectiveness of the intervention) to address the uncertainties around the ICERs.

In conclusion, we hypothesise that the additional upfront cost of delivering the intervention will be counterbalanced by improvements in clinical practice and patient related outcomes, thereby rendering the CARRS QI strategy cost-effective. The results of this study will be of immediate relevance for decision makers of all sorts –patients, healthcare providers, and policy makers– concerning implementation of
this healthcare delivery intervention to improve diabetes care goals.

Abbreviations
- €: Euros; BP: Blood pressure; CARRS: Centre for Cardiometabolic Risk Reduction in South Asia; CC: Care coordinator; CI: Confidence interval; CRF: Case report form; CVD: Cardiovascular disease; DBP: Diastolic blood pressure; DS-HER: Decision support electronic health records; DSS: Decision support software; ECG: Electrocardiogram; EQ-5D 3 L: European Quality of Life 5 dimensions - 3 levels; GDP: Gross domestic product; HbA1c: Glycated haemoglobin; ICER: Incremental cost-effectiveness ratio; LDLc: Low-density lipoprotein cholesterol; LMIC: Low- and middle-income countries; QALYs: Quality adjusted life years; QI: Quality improvement; SBP: Systolic blood pressure; UK: United Kingdom; UKPDS: United Kingdom Prospective Diabetes Study; USE: United States dollar.

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Availability of data and materials
Not applicable as this manuscript represent a study protocol for the cost-effectiveness analysis.

Authors’ contributions
KS, NT, DP, KMVN, MKA, and VP have formulated the design of the economic effectiveness analysis. KS, NT, MK, KMVN and DP have composed the questionnaires. All authors have taken part in preparation of the manuscript and have reviewed critically for intellectual content and approved the final version.

Ethics approval and consent to participate
Institutional ethics committees at each participating site and the research coordinating centres (Public Health Foundation of India and Emory University, USA) approved the study and all physicians and patients gave written informed consent prior to participating in this study.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1Public Health Foundation of India, Center of Excellence - Center for Cardio-metabolic Risk Reduction in South Asia, 4th Floor, Plot No. 47, Sector 44, Institutional Area, Gurgaon, Haryana 122 002, India. 2Rollins School of Public Health, Emory University, 1518 Clifton Road, Emory 30322, USA. 3Department of Endocrinology & Diabetes, Amrita Institute of Medical Sciences, Amrita Vishwa Vidyapeetham, Ponekkara P.O., Kochi, Kerala 682 041, India. 4Department of Endocrinology, TMN College & BYL Nair Charity Hospital, Dr. A. L. Nair Road, Mumbai Central, Mumbai, Maharashtra 400 008, India. 5MV Hospital for Diabetes & Diabetes Research Centre, No 4, West Madha Church Street, Royapuram, Chennai, Tamil Nadu 600 013, India. 6Bangalore Endocrinology & Diabetes Research Centre, #35, 5th Cross, Malleswaram Circle, Bangalore, Karnataka 560 003, India. 7Department of Endocrinology, St. John’s Medical College & Hospital, Sarjapur Road, Koramangala, Bangalore, Karnataka 560 034, India. 8Department of Endocrinology, Omosara General Hospital, 2nd Floor, Golden Jubilee Block, Azfalgunj, Hyderabad, Telangana 500 012, India. 9Department of Medicine, Section of Endocrinology and Diabetes, Aga Khan University, Stadium Road, Karachi 74800, Pakistan. 10Department of Endocrinology & Metabolism, All India Institute of Medical Sciences, Biotechnology Block, 3rd Floor, Anspi Nagar, New Delhi 110 029, India. 11Endocrine Unit, Department of Medicine, Goa Medical College, Bambolim, Goa 403202, India. 12Public Health Foundation of India, 4th Floor, Plot No. 47, Sector 44, Institutional Area, Gurgaon, Haryana 122 002, India. 13Rollins School of Public Health, Emory University, 1518 Clifton Road, Emory 30322, USA. 14Rollins School of Public Health, Emory University, 1518 Clifton Road, Atlanta, GA 30322, USA. 15Department of Endocrinology & Metabolism, All India Institute of Medical Sciences, Biotechnology Block, 3rd Floor, Rm #312, Anspi Nagar, New Delhi 110 029, India. 16Centre for Chronic Disease Control, C 1/52, Second floor, Safdarjung Development Area, New Delhi, 110016, India. 17London School of Hygiene and Tropical Medicine, Keppel St, Bloomsbury, London WC1E 7HT, UK.

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