A systematic review and meta-analysis of the impact of GLP-1 receptor agonists and SGLT-2 inhibitors on cardiovascular outcomes in biologically healthy older adults

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Abstract
Background: Unintentional weight loss is a hallmark of frailty and is associated with poor outcomes in older adults with type 2 diabetes. As such, the role of pharmacological therapies that facilitate weight loss – namely, sodium-glucose co-transporter-2 (SGLT-2) inhibitors and glucagon-like peptide-1 (GLP-1) receptor agonists – remains uncertain in fitter older adults. We performed a systematic review and meta-analysis to evaluate these agents on major adverse cardiovascular events (MACE) in older adults eligible for participation in cardiovascular outcome trials.

Methods: A literature search was performed in MEDLINE, EMBASE, CINAHL, Cochrane Central Registry of Controlled Trials (CENTRAL) and CNKI from inception to 29 June 2020. A class-specific meta-analysis was conducted in older adults (>65 years at recruitment) and compared with the similar analysis in younger (<65 years) adults.

Results: Of 761 unique studies identified, nine met the criteria for inclusion, five using GLP-1 receptor agonists and four with SGLT-2 inhibitors. GLP-1 receptor agonists in older adults were associated with a 15.3% (OR 0.847 (95% CI 0.788 to 0.910)) reduction in MACE events, similar to the 16% benefit seen in younger adults. The use of SGLT-2 inhibitors reduced MACE in older participants by 16.9% (OR 0.831 (95% CI 0.699 to 0.989)), numerically superior to the impact in younger patients (OR 0.936 (95% CI 0.787 to 1.113)).

Conclusions: GLP-1 receptor agonists and SGLT-2 inhibitors reduced MACE outcomes in older adults who were eligible to participate in clinical trials. Whereas this is reassuring for the biologically robust, it should not be extrapolated to frail older adults without further investigation.

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Key words: type 2 diabetes, older adults, major adverse cardiovascular events, cardiovascular outcome trials, GLP-1 receptor agonists, SGLT-2 inhibitors

Introduction
The International Diabetes Federation estimated the number of people living with diabetes worldwide at 463 million in 2019, 85% of whom had type 2 diabetes.1 This figure, however, does not include adults over the age of 80. In the UK, approximately 20% of the population over the age of 85 have diabetes and 27% of people in care homes, accounting for 15% of the British population living with diabetes.2

Frailty – that is, a reduction in physiological reserve with associated sarcopenia, weight loss and functional decline – is a recognised complication of diabetes. Older adults with frailty and diabetes are at a greater risk from medication side effects, polypharmacy and complications due to their co-morbidities.2 There has been a recent focus on addressing the over-treatment of frail older adults in order to reduce this risk after the publication of a national stakeholders document for the assessment and target setting,3 followed by the adoption of this standard by the National Institute for Health and Care Excellence (NICE; NG158 and NG160) and the General Medical Services Quality Outcome Framework (GMS QOF) for primary care. Within these guidance documents, the use of drugs that promote weight loss are discouraged as a potential risk for progression of frailty. Specifically, the use glucagon-like peptide-1 (GLP-1) receptor agonists and sodium-glucose co-transporter-2 (SGLT-2) inhibitors are highlighted as agents that may carry risk for older adults. It is important to remember, however, that the majority of older adults are not frail. This was recognised in the stakeholders guidance document, which recommended that biologically fit older adults living with diabetes should be treated in a similar manner to their chronologically younger counterparts.3 Notwithstanding this, there remain concerns among some healthcare workers that agents facilitating weight loss may augment progression to frailty.

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GLP-1 analogues mimic the incretin response to food, increasing insulin secretion, slowing gastric emptying and promoting satiety.\(^6\) In the younger population (\(< 65\) years of age) these are associated with approximately 4–6 kg weight loss.\(^5\) SGLT-2 inhibitors reduce glucose reabsorption in the proximal tubule of the nephron, thereby promoting calorie loss and producing a similar 4–6 kg weight loss.\(^6\) These medications have been proven to reduce the incidence of major adverse cardiac events (MACE; cardiovascular death, non-fatal myocardial infarction and non-fatal stroke) in high-risk trial populations with type 2 diabetes.\(^7\)

We planned to address the uncertainty as to whether these agents have similar cardiovascular benefits in older patients as is accepted in younger populations, or whether the impact on weight loss promotes frailty thereby failing to show benefit or even being associated with harm. We performed a systematic review and meta-analysis of cardiovascular outcome trials of GLP-1 receptor agonists and SGLT-2 inhibitors, where a priori or post hoc analyses were presented stratified by age.

**Methods**

The protocol is registered on the PROSPERO database (CRD42020200601) and is reported in accordance with the reporting guidance provided in the Preferred Reporting Items for Systematic Reviews and Meta-Analyses Protocols (PRISMA-P) statement.\(^8,9\)

**Eligibility criteria**

Studies were selected according to the following criteria:

- **Study design**: Only prospective randomised controlled trials were included. Retrospective comparative cohort studies, case-control or nested case-control studies and case series were excluded.
- **Participants**: Any study that included people with type 2 diabetes and presented an a priori or post hoc age-stratified analysis with an event rate reported for ‘older adults’. Any age \(\geq 65\) years was regarded as satisfactory for ‘older’.
- **Interventions**: Studies that used a GLP-1 receptor antagonist or SGLT-2 inhibitor as the agent of interest.
- **Comparators**: The comparator could be placebo or standard care management aiming for glycaemic equipoise.
- **Outcome measures**: Adjudicated MACE outcome was required. This could be a 3-point MACE (cardiovascular death, non-fatal myocardial infarction or non-fatal stroke), a 4-point MACE (3-point plus hospitalisation for acute coronary syndrome) or MACE+ (3- or 4-point MACE and hospitalisation for heart failure).

**Information sources and search strategy**

We searched the following electronic databases: MEDLINE, EMBASE, CINAHL, Cochrane Central Registry of Controlled Trials (CENTRAL) and CNKI from inception to 29 June 2020. Only English language manuscripts were included. The reference lists of included studies were also searched for additional studies.

**Screening, selection and data extraction procedure**

Both authors screened all articles identified from the search independently, starting with titles and abstracts and then full texts were examined in detail. Any disagreements would have been resolved by discussion; however, both authors reached consensus on first review.

**Risk of bias assessment**

All studies were assessed for risk of bias (ROB) using the Cochrane ROB 2.0 tool in five domains: bias arising from the randomisation process, due to deviations from intended interventions, due to missing outcome data, in the measurement of the outcome and in the selection of the reported result.\(^10\) The signalling questions were judged as yes, probably yes, probably no, no, or no information, resulting in an overall judgement of study ROB as high, uncertain or low.

**Data synthesis and additional analyses**

Study characteristics are presented in a summary of characteristics table. Pooled estimates using the random-effects model, which is a more conservative estimate of treatment effect, were calculated for all treatments combined, then stratified for SGLT-2 inhibitors and GLP-1 receptor agonists. All analyses were performed using OpenMeta software. Heterogeneity was quantified by estimating the variance between studies using the \(I^2\) statistic.

**Results**

After duplications were removed, our initial search identified a possible 761 publications. The initial screen of abstracts reduced this to 103 manuscripts which were reviewed in full (Figure 1). Nine studies met our inclusion criteria, five using GLP-1 receptor agonists and four using SGLT-2 inhibitors. Descriptive details of

![](https://example.com/figure1.png)

**Figure 1.** PRISMA flow chart
these studies are presented in Tables 1 and 2. One notable study that could not be incorporated was the ELIXA (Lixisenatide in Patients with Type 2 Diabetes and Acute Coronary Syndrome) trial which did not report any age-specific findings.11

### Glucagon-like peptide-1 (GLP-1) receptor agonists

There were five placebo randomised controlled trials of subcutaneously administered GLP-1 receptor agonists, all in addition to standard of care including optimisation of blood pressure and lipid control. Four were post-marketing cardiovascular safety studies that allowed escalation of non-incretin-based therapy in order to match glycaemic control (EXSCEL using exenatide,12 LEADER using liraglutide,13 HARMONY using albiglutide14 and REWIND using dulaglutide15) with a trial duration ranging from 1.8 to 5.4 years. SUSTAIN-6 testing semaglutide was a placebo-controlled pre-approval regulatory study with a shorter duration (2.0 to 5.0 years) to the meta-analysis.

Table 1 Placebo-controlled trials of glucagon-like peptide-1 receptor agonists that reported 3-point MACE outcomes in older adults

<table>
<thead>
<tr>
<th>Trial</th>
<th>Year</th>
<th>Number of participants</th>
<th>Drug</th>
<th>‘Older’ age</th>
<th>Drug events</th>
<th>Placebo events</th>
<th>Hazard ratio</th>
<th>95% CI</th>
<th>% established CV disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>LEADER</td>
<td>2016</td>
<td>9340</td>
<td>Liraglutide</td>
<td>60</td>
<td>468/3471</td>
<td>528/3548</td>
<td>0.90</td>
<td>0.79 to 1.02</td>
<td>81</td>
</tr>
<tr>
<td>SUSTAIN-6</td>
<td>2016</td>
<td>3297</td>
<td>Semaglutide</td>
<td>65</td>
<td>557/93</td>
<td>76/805</td>
<td>0.72</td>
<td>0.51 to 1.02</td>
<td>83</td>
</tr>
<tr>
<td>EXSCEL</td>
<td>2017</td>
<td>14752</td>
<td>Exenatide</td>
<td>65</td>
<td>426/2964</td>
<td>512/2975</td>
<td>0.80</td>
<td>0.71 to 0.91</td>
<td>73</td>
</tr>
<tr>
<td>Harmony</td>
<td>2018</td>
<td>9463</td>
<td>Albiglutide</td>
<td>65-74</td>
<td>154/1771</td>
<td>166/1838</td>
<td>0.97</td>
<td>0.78 to 1.21</td>
<td>100</td>
</tr>
<tr>
<td>REWIND</td>
<td>2019</td>
<td>9901</td>
<td>Dulaglutide</td>
<td>66</td>
<td>331/2314</td>
<td>384/2350</td>
<td>0.86</td>
<td>0.74 to 1.00</td>
<td>31</td>
</tr>
</tbody>
</table>

### Sodium-glucose co-transporter-2 (SGLT-2) inhibitors

Four studies met our a priori selection criteria; however, one study (The Canagliflozin and Cardiovascular and Renal Events in Type 2 Diabetes; The CANVAS program) reported their findings in events per 1,000 patient years rather than absolute numbers,
rendering it impossible to include these data in the meta-analysis. The findings from CANVAS are therefore presented in the summary in Table 2, but not in the meta-analysis (Figure 2B).

The remaining three studies explored the role of empagliflozin after an acute coronary event (EMPA-REG), dapagliflozin in those with established or at high risk of developing cardiovascular disease (DECLARE) and canagliflozin in renal impairment (estimated glomerular filtration rate (eGFR) 30–90 mL/min; CREDENCE). All studies were placebo controlled in addition to standard of care with the intention of achieving glycaemic equipoise and met their primary endpoint of cardiovascular safety. EMPA-REG and CREDENCE achieved superiority compared with placebo in the total population whereas, in DECLARE, dapagliflozin only achieved the latter of the two

Figure 2. Major adverse cardiovascular events (MACE) in placebo-controlled trials comparing (A) glucagon-like peptide-1 receptor agonists (GLP-1RA) and (B) sodium-glucose co-transporter-2 inhibitors (SGLT-2i) in addition to standard of care in older adults (>65 years).

Figure 3. Major adverse cardiovascular events (MACE) in placebo-controlled trials comparing (A) glucagon-like peptide-1 receptor agonists (GLP-1RA) and (B) sodium-glucose co-transporter-2 inhibitors (SGLT-2i) in addition to standard of care in older adults (>65 years).
endpoints from the split alpha between 3-point MACE and hospitalisations due to heart failure.

In combination, there were 13,091 older adults considered in the analysis of the SGLT-2 inhibitors (Figure 2B). The 749 events in the 7,051 individuals in the treatment group (10.62%) was superior to the 722 events in the 6,040 individuals in the placebo group (11.95%) by 17% (OR 0.831 (95% CI 0.699 to 0.989); test for heterogeneity I²=54.8%, p=0.11). This absolute risk reduction of 1.33% translates to a number needed to treat of 75 people for 3.5 years to prevent one cardiovascular event. This compares favourably to younger patients in whom there was a non-significant 0.2% absolute risk reduction (OR 0.936 (95% CI 0.787 to 1.113); Figure 3B). Of interest, the point estimates of CANVAS program for both older (OR 0.80 (95% CI 0.67 to 0.95)) and younger (OR 0.91 (95% CI 0.76 to 1.10)) participants was similar to the calculated odds ratio in the other studies, suggesting that, had they reported absolute numbers and been included in the meta-analysis, they would not have materially altered the findings.

**Discussion**

This systematic review has demonstrated the benefit of treatment with a GLP-1 receptor agonist or an SGLT-2 inhibitor in older adults who were eligible for inclusion in these large randomised controlled trials. In both cases the agents of interest were beneficial in older adults, with a smaller number of patients required to be treated in order to gain benefit. Indeed, in the case of SGLT-2 inhibitors, benefit was only seen in older adults whereas in younger populations no significant benefit was demonstrated. This is of particular relevance given the recent update to the UK guidance for the management of older adults with diabetes, which emphasises the need to assess frailty in individuals and subsequently individualise the treatment options.3 This stakeholder’s document highlighted the observation that weight loss in older adults has no proven benefit; indeed, it may exacerbate sarcopenia and thus have a detrimental impact on prognosis.21 As a result, the stakeholder’s position statement recommended caution in the use of agents that may promote weight loss. The current analysis suggests, however, that fitter older adults, who would have been eligible for inclusion in clinical trials, would benefit from treatment with SGLT-2 inhibitors or GLP-1 analogues.

It is important to acknowledge, however, that older clinical trial participants may not be truly representative of the general population. Indeed, the term ‘older adults’ can span more than 35 years, from 65 years to centenarians, with some being robust in employment and independently active whilst others are in the terminal decline of multi-morbidity and frailty. The physiological differences attributable to frailty are substantial; however, current treatment strategies are based on extrapolation from the outcomes in chronologically matched patients.22 Frail older adults are rarely included in clinical trials, with exclusion criteria to recruitment such as polypharmacy, cognitive decline and multiple co-morbidities being common.23 For this reason, older adults included in cardiovascular outcome trials tend to be biologically healthier than the general older population.24 Even within these populations, trials of glycaemic control in older adults highlight an exaggerated ‘U-shaped’ curve with higher mortality at both low and higher HbA₁c.25,26 To date there are very limited data on the optimal medication choices for frail older adults, and fewer still on the individualisation of care in those living with frailty.27 There is, however, a pressing need to explore such populations, given the significant impact that heart failure and stroke have on functional capacity in older adults. These complications of diabetes are significant contributors to frailty progression. The co-existence of heart failure and diabetes has a poorer prognosis than many cancers.28 An anticipated life expectancy of less than 7 years is a common exclusion criterion,29 which would thereby exclude the patients with the most to gain in our analysis from the studies to evaluate their efficacy.

This systematic review has highlighted the importance and potential benefits of using both GLP-1 agonists and SGLT-2 inhibitors to reduce cardiovascular outcomes in older adults with type 2 diabetes. A key limitation is that only eight studies were eligible for inclusion. Therefore, it is very important that further research in this field is undertaken. It is also important to acknowledge that 3-point MACE does not represent the only outcome of interest for older adults living with diabetes. It is accepted that SGLT-2 inhibitors, as a class, improve outcomes for people living with heart failure. Hospitalisation from heart failure is important for older adults with diabetes; however, these outcomes are not stratified by age and thus were not included in this meta-analysis. In a population in whom weight loss is associated with frailty and poor cardiovascular outcomes, the absence of any adverse cardiovascular signals should be reassuring to use SGLT-2 inhibitors in older adults with heart failure.
Conclusions
On the basis of these data, we would advocate the use of GLP-1 analogues and SGLT-2 inhibitors in any older adult with mild or moderate frailty, particularly when the frailty is a function of cardiovascular disease itself. Further research is required to determine whether the benefits are extended to older adults with severe frailty or patients at lower risk of cardiovascular events.

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Compliance with ethics guidelines This was a retrospective meta-analysis of published data. All studies incorporated in the analysis had been through appropriate ethics review and approvals.

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