Suitable Use of Injectable Agents to Overcome Hypoglycemia Risk, Barriers, and Clinical Inertia in Community-Dwelling Older Adults with Type 2 Diabetes Mellitus

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Abstract
The management of type 2 diabetes mellitus in older adults requires a comprehensive understanding of the relationship between the disease (medical) and the functional, psychological/cognitive, and social geriatric domains, to individualize both glycemic targets and therapeutic approaches. Prevention of hypoglycemia is a major priority that should be addressed as soon as its presence or risk is detected, adjusting the target and therapeutics accordingly. Nonetheless, treatment intensification should not be neglected when applicable, consistent with recommendations from organizations such as the American Geriatrics Society and the American Diabetes Association, to reduce not only long-term macrovascular and microvascular complications (individualization), but also short-term complications from hyperglycemia (polyuria, volume depletion, urinary incontinence). Such complications can negatively impact the physical and cognitive function of older adults, worsen their quality of life, and additionally affect their families and society. We emphasize individualization, utilizing the multiple classes of antihyperglycemic agents available. Metformin remains as first-line therapy, and additional agents offer advantages and disadvantages that ought to be considered when developing a patient-centric plan of care. For selected cases, injectable therapies such as long-acting basal insulin analogs and glucagon-like peptide-1 receptor agonists can offer advantages to counter hypoglycemia risk, patient-related barriers, and clinical inertia. Furthermore, some injectable agents could potentially simplify regimens while providing safe and effective glycemic control. In this review, we discuss the use of injectable therapies for selected community-dwelling older adults, barriers to transition to injectable therapy, and measures aimed at removing these barriers and assisting physicians and their teams to transition older patients to injectable therapies when appropriate.

1 Introduction
Type 2 diabetes mellitus (T2D) is a chronic, progressive disease that can lead to multiple macrovascular and microvascular complications. It affects individuals, their families (time, non-paid caregiver costs, formal support), and society (healthcare costs and utilization, resources, and policies) [1]. The clinical course of T2D in older adults is heterogeneous and complex, impacted by age-related diseases (e.g., obesity, depression, and geriatric syndromes) and aging itself [2]. Successful and safe management of a patient with impaired physical or cognitive function will greatly rely on factors such as social support and resources [1–3]. Periodic monitoring and adjustment of targets and pharmacologic strategies require understanding the four geriatric domains in the older adult—psychological, medical, social, and functional—and optimizing targets and therapies in a timely fashion, while preventing complications (from hyperglycemia, hypoglycemia, or glycemic variability) [1–4]. Thus, major organizations highlight the need for treatment individualization, with special focus on the older population, incorporating geriatric assessments for function, cognition, and the burden of complications and comorbidities [5–10]. The common objective is to implement strategies that match the special needs of the aging population with diabetes [1–10].
The therapeutic options to accomplish safe glycemic control are multiple, from oral monotherapy to more complex regimens, including multiple options for injectable agents [9]. Recent changes to updated guideline recommendations emphasize the consideration of cardiovascular benefits into the decision making for antihyperglycemic agents [9, 10]. The evidence supports the use of oral agents such as sodium–glucose co-transporter 2 (SGLT2) inhibitor among patients with T2D with co-existent coronary artery disease and heart failure [9, 10]. Alternatively, among patients in need of treatment modification, the use of injectable agents can be hindered by fear of side effects such as weight gain and hypoglycemia, as well as concerns related to treatment burden (costs and complexity) and negative medication beliefs [11, 12], and fear of needles [13]. Also, in our clinical experience, patients often oppose these agents due to feelings of disappointment and a sense of defeat by the disease itself. These barriers, as well as clinical inertia, lead to inadequate glycemic control [13].

Thus, the scope of this paper is to review the value of alternative therapeutic approaches such as those offered by injectable antihyperglycemic agents among community-dwelling older adults with T2D who, upon target individualization, remain uncontrolled, have high hypoglycemia risk on their established regimen, or for whom short-acting insulin is not applicable due to the complexity of the regimen. This review does not address the hospital, nursing home, or end-of-life level-of-care settings. The literature search focused on peer-reviewed publications available in PubMed and EMBASE, using key words that connected the above-mentioned topics, using generic search terms (e.g., “injectable therapies in older adults with T2D”) and more specific terms (e.g., the name of each injectable agent currently available, with search limits in older adults or aging population). The search limits included publications in the past 5 years, aiming to compile the newest reports on pharmacologic agents, guidelines, and opinions about their use in older adults. This review is intended for general practitioners and specialists caring for older adults with T2D, for whom treatment intensification is not contraindicated, but actually required, to facilitate discussions with patients and their caregivers, and to reduce hypoglycemia and treatment burden, especially considering that some newer injectable agents can offer the advantage of weekly regimens.

2 Managing Type 2 Diabetes (T2D) in Older Adults

The approach to treating an older adult with diabetes requires understanding that, on the one hand, factors specific to the disease and the aging patient affect diabetic pathophysiology [14], and, on the other hand, the four geriatric domains are intertwined and affect diabetes self-management, self-efficacy, adherence, and the risk of medication errors and hypoglycemia (Fig. 1) [1–10]. Furthermore, the plan of care should incorporate functional status and quality of life, reducing, when possible, medicine burden, and actively identifying risks [15]. More recently, the need for a comprehensive geriatric assessment has been highlighted by an understanding of the role of frailty in diabetes and individualization of targets and strategies [16, 17]. Hypoglycemia risk is high, even in those with poor glycemic control [18], and it must be viewed as a serious adverse event [19]. Advancing age is the most common risk factor for hypoglycemia-related hospitalizations [20]. In parallel, diabetes is associated with impaired physical function (functional disability, frailty, and accelerated muscle loss), psychological function (depression, cognitive impairment, and dementia) [1–10], and social function (high disease-related burden, costs, and economic distress) [21, 22]. The management starts by individualizing targets and pharmacologic approaches based on disease duration, multimorbidity, hypoglycemia risk, feasibility and support, treatment costs, quality of life, and life expectancy [1–10].

2.1 Adopting a Healthy Lifestyle and Environment

Lifestyle modification is the foundation of T2D management. Strategies include (1) setting goals based on patients’ input, language, numeracy, and cultural barriers; (2) integrating evidence-based guidelines and clinical information tools; and (3) encouraging connection with community resources, such as activity groups [3].

Intentional weight loss is not desirable in a 68-year-old patient with newly diagnosed cancer, a 75-year-old patient with dementia, or an 83-year-old patient with disability and dependence. However, older adults with T2D and obesity, who also have preserved physical and cognitive function,
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can benefit from modest intentional weight loss as part of their plan of care [23]. Proper nutrition, physical activity, and the four types of recommended exercise can be feasible, effective, and successfully implemented [23]. Education is paramount, addressing the need to monitor and adjust medications to avoid hypoglycemia.

### 2.2 Pharmacologic Treatment Beyond Metformin and Other Oral Antihyperglycemic Agents

Metformin remains the first-line treatment for T2D [3, 9, 10]. However, diabetes is chronic and progressive, with altered pathophysiology, as mentioned above. As a consequence, older adults are likely to require additional therapies to achieve the desired individualized target. The newest guidelines of the American Diabetes Association [9] recommend a step-wise approach, considering first the presence of established atherosclerotic cardiovascular disease (ASCVD), and implementation of SGLT2 inhibitors, or glucagon-like peptide 1 receptor agonists (GLP-1 RAs), because of their demonstrated cardiovascular disease benefit [9]. When ASCVD is not present, then the next big factor to consider is the presence or risk of hypoglycemia. Upon treatment individualization and implementation of the appropriate therapies, some older patients may achieve control, while others may need further treatment intensification to reach their individualized targets [1–10]. The subsequent selection of antihyperglycemic agents should be based on the pharmacologic profile and safety of the medication. Our review does not focus on T2D management in long-term care and skilled nursing facilities, for which separate guidelines are available [24]. Instead, we highlight the scenarios of community-dwelling older adults.

Fig. 1 A comprehensive approach to diabetes management in older adults. Evaluation and understanding of the four geriatric domains are fundamental for the implementation of best practices in older adults with diabetes. The four domains are intertwined, with interactions that impact diabetes self-management and self-efficacy, adherence, and risks. A healthy older adult can receive aggressive interventions in the setting of excellent self-management and self-efficacy skills and access to care and social support. As the person ages, not only does T2D run its natural course of progressive disease, but pharmacologic management becomes more challenging because of multimorbidity, functional impairment, and cognitive decline, among others, which limit self-management and self-efficacy. Diabetes targets need to be reviewed and adjusted accordingly. Avoiding hypoglycemia is the priority, while preserving quality of life is a major outcome and marker of successful management. As life expectancy decreases, therapy will focus on the prevention of hyperglycemia emergencies, diabetic polyuria, and dehydration. T2D type 2 diabetes. Adapted from Valencia and Florez [1]
who, upon target individualization and therapy with metformin and the next-line agents, may still require treatment adjustments, individualized to their clinical presentation. 

The most commonly used oral antihyperglycemic agents are summarized in Table 1 [1–3, 5–10, 25, 26].

### 2.3 Injectable Non-insulin Agents

GLP-1 RAs may be advantageous, as they have shown beneficial effects on body weight and are associated with a low risk of hypoglycemia in clinical trials [27].

Recently, significant attention has been placed on addressing potential additional cardiovascular benefits. While positive outcomes were observed in cardiovascular outcome studies with liraglutide (LEADER) [28], semaglutide (SUSTAIN) [29], and dulaglutide (REWIND) [30], the case was not the same with lixisenatide (ELIXA) [31] or exenatide (EXSCEL) [32]. A recent review focusing on cardiovascular outcomes of all GLP-1 RAs [33] summarized the available evidence on cardiovascular outcomes, and as not every agent has proven cardiovascular benefit, these benefits may not be a class effect. Moreover, there are observable differences in

| Table 1 Oral pharmacologic options for older patients with type 2 diabetes |
|-----------------------------|---------------------------|---------------------|---------------------|
| **Class**                   | **Agents**                | **Advantages**      | **Disadvantages**   |
| Biguanide [1–3, 5–10]       | Metformin                 | Low cost            | Lactic acidosis risk (rare) |
|                             |                           | Improvement in dyslipidemia | Contraindicated when patient has renal insufficiency or significant heart failure |
|                             |                           | Proven safety and effectiveness | GI side effects |
|                             |                           | No hypoglycemia       | Risk of vitamin B12 deficiency |
|                             |                           | Lowers CV risk        | May cause weight loss or GI side effects in frail patients |
|                             |                           | Lower cancer risk than other therapies | |
| SU [1–3, 5–10]              | Glimepiride, glipizide   | Low cost             | Weight gain         |
|                             |                           | Extensive experience  | Hypoglycemia        |
|                             |                           | Lowers microvascular risk | Caution recommended in patients with renal, cardiac, or hepatic insufficiency |
|                             |                           | Better CV safety than other oral agents | Possible increased risk of CVD and CV mortality |
| GLSi [1–3, 5–10]            | Acarbose, miglitol        | Lower hypoglycemia and weight gain than SU | GI side effects (require gradual titration) |
|                             |                           | Better CV safety than other oral agents | Potential hepatotoxicity with acarbose |
|                             |                           | Reduces PPG excursions | Miglitol contraindicated in people with renal failure |
| TZD [1–3, 5–10]             | Pioglitazone, rosiglitazone | Low cost            | Moderate A1C lowering |
|                             |                           | Low hypoglycemia risk | Frequent dosing schedule |
|                             |                           | Well tolerated and effective | Hepatic toxicity reported |
|                             |                           | Can be used in renal impairment | Weight gain due to fluid retention |
|                             |                           | Low cost             | Increased risk of bone loss and fractures |
| DPP-4i [1–3, 5–10]          | Sitagliptin, vildagliptin, saxagliptin, linagliptin, alogliptin | Low hypoglycemia risk | Edema/heart failure |
|                             |                           | Well tolerated       | Possible increased CV risk |
|                             |                           | Weight neutral       |                          |
|                             |                           | May help preserve β cells |                          |
| SGLT2i [1–3, 5–10, 25, 26]  | Canagliflozin, dapagliflozin, empagliflozin, ertugliflozin | Low hypoglycemia risk | High cost |
|                             |                           | Weight loss          | GU infections |
|                             |                           | Lower systolic blood pressure | Volume depletion, hypotension, and dizziness |
|                             |                           | Well tolerated       | Increased LDL-C |
|                             |                           | Efficacious and safe in older adults with chronic kidney disease | Concerns regarding long-term impact on CV risk and carcinogenicity |
|                             |                           | Associated with lower CVD event rate and mortality in patients with CVD | |

Oral agents are beyond the scope of this review but are included here for easy reference:

- A1C: glycated hemoglobin
- CV: cardiovascular
- CVD: cardiovascular disease
- DPP-4i: dipeptidyl peptidase 4 inhibitors
- GI: gastrointestinal
- GLSi: glucosidase inhibitors
- GU: genitourinary
- LDL-C: low-density lipoprotein cholesterol
- PPG: postprandial glucose
- SGLT2i: sodium–glucose co-transporter 2 inhibitor
- SU: sulfonylureas
- TZD: thiazolidinediones

△ Adis
the extent of the cardiovascular benefits between the GLP-1 RAs with positive outcomes.

On the other hand, while most studies did include older patients, few focused specifically on the older age group. The GetGoal-O study recruited patients aged ≥70 years and randomized them to lixisenatide or placebo [34]. The results showed efficacy in reducing glycated hemoglobin (A1C), postprandial plasma glucose, and body weight. However, patients experienced more frequent nausea, vomiting, and diarrhea. Similarly, a prior study with liraglutide found that older age was associated with more gastrointestinal side effects [35]. Nevertheless, studies with liraglutide [36] and lixisenatide [37] have shown GLP-1 RAs can be well tolerated by older adults. Proper patient education and monitoring can enhance the initial implementation, and slow titration should be considered, particularly in this age group.

A 2019 cost-effectiveness analysis using the Swedish Institute of Health Economics (IHE) Diabetes Cohort Model compared different types of GLP-1 RAs with each other and with insulin [38]. The design was modeled on T2D patients who did not achieve control on metformin or basal insulin, and results favored the once-weekly formulation of semaglutide over lixisenatide and dulaglutide.

### 2.4 Insulin

Diabetes is a chronic progressive disease, and many older patients will eventually require, and benefit from, insulin therapy. We recommend educating patients about this fact earlier in their disease, where we observed less reluctance to initiate injectable therapies. The following is a review of studies addressing multiple insulin therapies.

In the Treat-to-Target trial, 756 patients, mean age 55 years, were randomly assigned to GLA-100 or neutral protamine Hagedorn (NPH) insulin. While both treatments were effective, those on GLA-100 experienced fewer documented hypoglycemia events [39]. More recently, a pooled analysis of five randomized clinical trials (RCTs) compared the safety of GLA-100 with NPH [40]. Regarding the subset of older adults (329 treated with GLA-100, 275 treated with NPH), greater event rates of hypoglycemia occurred in those receiving NPH, albeit without statistical significance. Notably, patients on NPH were receiving a once-daily dosage, which is not the proper practice based on the drug’s half-life. A pooled analysis of 675 older patients with T2D found that patients treated with GLA-100 experienced better glycemic control and reduced incidence of hypoglycemia when compared with sulfonylureas, NPH, NPH 30/70, and insulin lispro (iLis) mix 75/25 [41].

With regard to fixed insulin combinations, the DURABLE trial was a 30-month, multicenter RCT that randomized 258 patients to insulin lispro (iLis) mix 75/25 (intermediate- and short-acting iLis) and 222 patients to GLA-100 [42–45]. After 24 weeks, iLis resulted in slightly lower HbA1c levels and slightly higher reductions of patients achieving target HbA1c < 7.0% (53 mmol/mol), though these patients also experienced more weight gain and higher rates of overall hypoglycemia, but lower rates of nocturnal hypoglycemia [45]. While intermediate-acting preparations have been used as basal insulin, they require at least two injections per day and have greater hypoglycemia risk. We recommend avoiding them, except in the setting of economic limitations and lack of access to other injectable preparations (see Table 2) [1–3, 5–10, 46, 47].

Newer long-acting insulins, such as insulin glargine 300 units/mL (GLA-300) and insulin degludec (IDeg) 100 units/mL and 200 units/mL, have been shown to provide a more constant pharmacokinetic profile with less hypoglycemia risk [48–50]. EDITION 3 was a multicenter, open-label, parallel-group study that randomized 878 patients (age 57.7 ± 10.1 years, disease duration 9.8 ± 6.4 years) to GLA-300 or GLA-100 once daily for 6 months. The researchers found similar A1C reduction in both groups (change from baseline was 0.04% [95% confidence interval (CI) − 0.09 to 0.17] or 0.4 mmol/mol [−1.0 to 1.9]) and lower hypoglycemia risk with GLA-300 (relative risk [RR] reduction of 24%; RR 0.76, 95% CI 0.59–0.99) [48]. A post hoc, patient-level meta-analysis of data from EDITION 2 (n = 811) and EDITION 3 (n = 878) showed that patients treated with GLA-300 had comparable glycemic control when compared with those treated with GLA-100, with reduced confirmed or severe nocturnal hypoglycemia (RR 0.64, 95% CI 0.48–0.85) and confirmed or severe hypoglycemia at any time of day (RR 0.77, 95% CI 0.65–0.91) [51]. Similar results were observed in another post hoc meta-analysis, which included patients from EDITION 1, 2, and 3 and evaluated outcomes over a 12-month follow-up period [52]. Nocturnal hypoglycemia risk was lower with GLA-300 (RR 0.85, 95% CI 0.77–0.92), and glycemic control defined as A1C < 7.0% without nocturnal hypoglycemia was achieved by 24% more patients with GLA-300 than with GLA-100 (RR 1.24, 95% CI 1.03–1.50). Data from clinical trials is supported by real-world evidence utilizing electronic medical records. DELIVER Naïve compared insulin-naïve patients initiating GLA-300 and GLA-100. After 6 months’ follow-up, patients treated with GLA-300 had significantly greater reductions in A1C (− 0.52 vs − 1.30; p = 0.003), with more GLA-300-treated patients achieving A1C < 7.0% (21.9% vs 17.4%; p = 0.003) without hypoglycemia [53]. In DELIVER 3, which studied older patients (≥65 years) with T2D switching to GLA-300 from first-generation basal insulins (GLA-100 or insulin detemir), A1C reductions were greater/similar with GLA-300, and A1C goal achievement was similar in both cohorts, but GLA-300-treated patients generally had less hypoglycemia (event rate: adjusted rate ratio 0.63, 95% CI 0.53–0.75; p < 0.001) [54]. Post-hoc and
real-world analyses offer important insight, especially when addressing the outcomes in the subpopulation of older adults [52]. The SENIOR study was the first prospectively designed study to address the efficacy and safety of insulin glargine in older people (age ≥ 65 years) with T2D [55]. With a multinational, multicenter, randomized, open-label study design, 1014 patients were allocated to Gla-300 or Gla-100. While glycemic control was similar, the events of confirmed hypoglycemia were low and similar in both groups, with differences in the subgroup age ≥ 75 years. In these patients, there were fewer events with Gla-300 than with Gla-100 (1.12 vs 2.71, rate ratio 0.45, 95% CI 0.25–0.83). In parallel, the DEVOTE study randomly assigned 3818 patients to IDeg and 3819 patients to Gla-100, to assess cardiovascular safety [56]. The primary outcome (a composite of first occurrence of an adjudicated major cardiovascular event) occurred in 325 patients (8.5%) with IDeg and in 356 (9.3%) with Gla-100 (HR 0.91, 95% CI 0.78–1.06; p < 0.001 for non-inferiority), thus concluding IDeg was non-inferior to Gla-100 with respect to cardiovascular safety. Real-world data has shown that patients switching to IDeg from other basal insulins had significantly improved A1C (0.58%; p < 0.001), without significant weight gain [57].

On the other hand, studies addressing long-acting basal insulins could offer potential applications, especially for the older population already facing polypharmacy or limitations to administration of injectable agents. A small pilot study in Japan found a thrice-weekly IDeg regimen was well tolerated in older adults with poorly controlled diabetes who had difficulty performing self-injection of insulin [58]. Using continuous glucose monitoring to assess and compare glycemic control, there were no differences between thrice-weekly versus once-daily injection groups for glucose < 70 mg/dL (1.3 ± 2.5% vs 2.4 ± 3.1%; p = 0.39) or glucose > 200 mg/dL (15.6 ± 18.0% vs 7.2 ± 12.1%; p = 0.22) [58]. Future studies confirming these results would support strategies where home-health nursing services (or family members, if well trained) could administer ultra-long-acting insulin formulations to elderly patients who cannot complete this task, thus becoming a good practical treatment option for these patients.

2.5 Case Studies for Whom Injectable Agents May Be Considered for the Plan of Care

Case Study 1: A 72-year-old female, with obesity, metabolic syndrome, diabetic microvascular complications, but

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### Table 2 Injectable pharmacologic options for older patients with type 2 diabetes

| Class                     | Agents                                                                 | Advantages                                                                 | Disadvantages                                                                 |
|---------------------------|------------------------------------------------------------------------|---------------------------------------------------------------------------|-------------------------------------------------------------------------------|
| GLP-1 RA [1–3, 5–10]     | Exenatide, liraglutide, dulaglutide, lixisenatide                      | Low hypoglycemia risk                                                     | High cost                                                                     |
|                           |                                                                        | Weight loss                                                                |                                                                               |
|                           |                                                                        | Reduced PPG excursions                                                     |                                                                               |
|                           |                                                                        | Reduces some CV risk factors                                               |                                                                               |
|                           |                                                                        | Once-daily and once-weekly formulations                                   |                                                                               |
| Intermediate-acting insulin [1–3, 5–10] | NPH | Long-term experience of use                                              | Significantly elevated risk of hypoglycemia                                 |
|                           |                                                                        | Nearly universal response                                                  |                                                                               |
| Long-acting insulin [1–3, 5–10] | Insulin glargine 100 units/mL, insulin glargine 300 units/mL, insulin detemir, insulin degludec (100 or 200 units/mL) | Provides relatively uniform insulin throughout the day                     | Hypoglycemia                                                                 |
|                           |                                                                        | Nearly universal response                                                  | Weight gain                                                                   |
|                           |                                                                        | Theoretically unlimited efficacy                                           |                                                                               |
|                           |                                                                        | Lower hypoglycemia risk than NPH                                           |                                                                               |
| Premix insulin [9, 10]    | 70/30 (NPH + regular, NPH + aspart), 75/25 (lispro protamine + lispro) | Reduced number of injections                                               | Hypoglycemia                                                                 |
|                           |                                                                        | Long-term experience of use                                                | Weight gain                                                                   |
| Fixed-ratio combination GLP-1 RA plus insulin [46, 47] | iGlarLixi (insulin glargine 100 units/mL + lixisenatide), IDegLira (insulin degludec 100 units/mL + liraglutide) | Reduced number of injections                                               | Moderate hypoglycemia risk                                                   |
|                           |                                                                        | Possible better glucose control and increased safety compared with         | Maximal doses                                                                |
|                           |                                                                        | individual components                                                      |                                                                               |

Short-acting synthetic insulin formulations are not mentioned as these are not usually started at the primary-care level because they require more specialized monitoring and titration, usually by an endocrinologist. Nonetheless, we emphasize that twice-daily regimens with combined insulins require an additional injection of short-acting insulin to cover lunch.

CV cardiovascular, GI gastrointestinal, GLP-1 RA glucagon-like peptide-1 receptor agonist, NPH neutral protamine Hagedorn, PPG postprandial glucose.
without known ASCVD, whose parents are still alive (aged 98 and 97 years). Her antihyperglycemic regimen involves maximum-dose metformin and a sulfonylurea. Her most recent glycated hemoglobin was 11%. She has a prior history of gallbladder disease and pancreatitis, and dipeptidyl peptidase 4 inhibitors and GLP-1 RAs are not an option for her. Adding a SGLT2 inhibitor as an oral agent can be an option, but due to the degree of elevation of A1C, she needs to be started on long-acting insulin, to be titrated by fasting glucose levels, while improving lifestyle to enhance postprandial control, with reassessment in 3 months to evaluate the need to add another agent.

Case Study 2: A 79-year-old male, with obesity, metabolic syndrome, microvascular and macrovascular complications, including ASCVD and heart failure, who lives independently at home, with only minor cognitive decline and no functional impairments, no falls, and no prior hypoglycemic events. His regimen includes metformin, an SGLT2 inhibitor, and basal insulin (0.9 units/kg/day). However, his most recent A1C was 9%, and fasting glucose levels averaged 110 mg/dL. Therefore, adding more basal insulin is not going to offer postprandial control. The patient can benefit from multiple options, including both oral agents and injectable agents. Treatment individualization requires understanding the costs and accessibility, and the patient may consider a cheaper medication, such as a sulfonylurea, but needs to monitor weight and hypoglycemia risk. At the 3-month follow up, A1C had improved to 8% (which is on target). However, the patient reported four events of hypoglycemia per week (despite being told to report to the team, the patient misunderstood and thought the hypoglycemic events were to be expected. Luckily, there were no adverse events from the hypoglycemia, since they all occurred at home). The team evaluated the alternatives that would offer similar glycemic success, but with less risk of hypoglycemia.

3 Barriers to Appropriate Transition to Injectable Antihyperglycemic Therapies

Initiating and scaling injectable therapies in older adults present a variety of challenges. A recent systematic review identified three main themes as barriers [59]: (1) individual—fear of pain and injections, and concerns about the side effects, including hypoglycemia; (2) healthcare professional—poor knowledge and skills, clinical inertia, fear of hypoglycemia, and language barriers; (3) system-related barriers—not having enough time to manage dose adjustments and monitor potential side effects, and lack of educational resources.

3.1 Hypoglycemia Risk and Fear of Hypoglycemia

Older adults with diabetes are at increased risk of hypoglycemia [1–4, 9, 11, 20]. Hypoglycemia-related hospital admissions are nearly two-fold higher for patients aged ≥75 years compared with those aged 65–74 years [60]. Consequences include dysrhythmias [61] (especially in those with cardiomyopathy) and accidents, falls, and related fractures [62]. Undoubtedly, preventing and minimizing hypoglycemia risk is one of the most important factors for determining glycemic goals and therapeutic approach [3, 9]. However, misunderstanding this approach by not increasing therapies to reach targets leads to clinical inertia.

3.2 Patient-Related Barriers

Personal barriers include perceived loss of personal control over treatment, the feeling of failure to control diabetes with oral agents, decreased flexibility, and increased inconvenience [9]. From a social perspective, negative reactions to insulin use from other people, the stigma and discrimination associated with the use of needles, and interference with social and work activities have been identified [59]. Finally, financial barriers can be due to increased diabetes healthcare costs, especially as older adults present multimorbidity and polypharmacy [1–3, 5–10].

3.3 Clinical Inertia: Delayed Initiation of Appropriate Therapy

Clinical inertia is a complex, multifactorial problem resulting from patient and physician barriers to treatment intensification [63]. A recent retrospective study from a US managed-care claims database of patients with uncontrolled T2D found that up to 72.8% of these patients experienced clinical inertia [64]. Of these, only up to 6.2% intensified treatment with insulin. Remarkably, a key barrier was older age itself. Clinical inertia has important health care consequences. A 1-year delay in insulin initiation, in conjunction with poor glycemic control, significantly increases the risk of T2D complications, including myocardial infarction, heart failure, and stroke [65]. A retrospective cohort study of patients with T2D, grouped per clinical inertia (failure to initiate insulin within 3 months of an A1C level ≥9.0% (75 mmol/mol) despite taking two oral antidiabetic drugs), found that clinical inertia was associated with a significantly shorter median time to progression of diabetic retinopathy (p = 0.02) and a higher incidence of diabetic retinopathy progression (10 vs 2.2 cases per 1000 person-months; p = 0.003) [66]. Receiving treatment from a general practitioner was the strongest risk factor for clinical inertia. Another retrospective study reported that in patients aged ≥65 years with T2D,
mean time of exposure above A1C 7.0% (53 mmol/mol) was 15 months [67].

A 2014 study evaluated the implementation of injectable agents between 2000 and 2009 in 51,771 patients with T2D who were receiving two oral agents and were followed for 2 years [68]. The majority (79.3%) was started on a third oral agent compared with those started on insulin (13.3%) or a GLP-1 RA (7.4%). Those started on insulin had greater improvements in glycemic control.

A 2017 analysis of > 10,000 individuals from a US clinical practice found that patients who were older (adjusted odds ratio [OR] 0.975, 95% CI 0.971–0.979) and had higher A1C values (OR 0.741, 95% CI 0.721–0.761) and Diabetes Complications Severity Index scores (OR 0.870, 95% CI 0.848–0.892) were significantly less likely to be prescribed a GLP-1 RA compared with basal insulin [69]. While we did not find any European study of similar design, a 2017 report from a very large database (> 400,000 patients) from Italy, The Netherlands, Spain, and the UK, described injectables as being used mostly as third-line agents [70]. The main factors driving treatment choice at any stage of intensification were older age, A1C, body mass index, renal and cardiac morbidity, and treatment history.

4 Transition to Injectable Agents to Overcome Hypoglycemia Risk, Patient-Related Barriers, and Clinical Inertia

Once basal insulin is indicated as the next-best step for intervention, the provider must first address hypoglycemia risk and patient-related barriers. Notably, as we describe in the upcoming section, many newer injectable agents offer reduced hypoglycemia risk and can effectively help the patient to reach the individualized target and accomplish glycemic control. We present a summary in Table 3 [1–10, 59–67].

4.1 Reducing Hypoglycemia Risk

Hypoglycemia events require a clear understanding of their etiology to avoid a recurrence. Details in the history may reveal that the patient accidentally injected the correct dose twice because of forgetting an earlier dose, or that the patient was interrupted during a meal that remained unfinished. In both scenarios, the regimen may remain effective and safe if the events are isolated and conditions do not change. Recurrent events can be a sign of cognitive decline or early self-care deficits. Glycemic targets need to be adjusted, and further coordination of services (formal or informal) will be required in order to deliver the injectable therapeutic plan and to avoid hypoglycemia.

Despite concerns regarding hypoglycemia risk, several studies have demonstrated that many injectable therapies can be safe when properly implemented. Beyond the known low risk of hypoglycemia with the most commonly used long-acting basal insulins (Gla-100 and detemir), newer formulations include more concentrated and longer-acting basal insulins. The analyses from the EDITION studies demonstrated a significantly lower risk of hypoglycemia with Gla-300, with fewer participants reporting nocturnal or severe hypoglycemia episodes, in both insulin-naïve [48] and insulin-experienced [49] patients. These findings were corroborated in the SENIOR study [55]. Similarly, IDeg has demonstrated reduced rates of nocturnal severe hypoglycemia compared with Gla-100 in patients with T2D [56].

Regarding the time of administration, data from two phase III, 26-week RCTs that compared IDeg with daily Gla-100 showed hypoglycemia episodes were greater when IDeg was administered at night rather than in the morning [71]. Basal insulin is usually administered at night to adjust dosage based on the next day’s first fasting serum glucose, but switching to morning administration, ideally after a steady effective dose has been established, may decrease events. The principle of “start low and go slow” does not preclude, but rather facilitates the process to further titrate medications to accomplish the individualized target goals.

On the other hand, GLP-1 RAs offer options from twice-daily (exenatide) to once-weekly (semaglutide, exenatide) treatment. A recent systematic review indicates that longer-duration GLP-1 RAs, such as once-weekly exenatide, may have less hypoglycemia risk than short-acting exenatide [72].

While the combination of injectable agents (e.g., basal insulin with pre-meal insulin, basal insulin with GLP-1 RAs) may fall under the specialist care of an endocrinologist, newer options are becoming available, with fixed dosages that could feasibly be implemented at the primary-care level. For example, short-acting GLP-1 RAs added to basal insulin resulted in a reduced number of hypoglycemia events when compared with the addition of short-acting insulin [73]. In this study, a subset of subjects were older. On the other hand, two fixed-ratio combinations of basal insulin plus GLP-1 RA have been approved and have demonstrated potential advantages. iGlarLixi (a combination of insulin glargine 100 units/mL and lixisenatide once daily) demonstrated greater glycemic control compared with the individual components, with no increased risk of hypoglycemia [74]. IDegLira (a combination of IDeg and liraglutide once daily) achieved glycemic control superior to that of IDeg at equivalent insulin doses, without an increased risk of hypoglycemia and with the benefit of weight loss [75].
Table 3  Strategies to overcome hypoglycemia risk, patient-related barriers, and clinical inertia in older patients

| Issues | Recommended strategies | Implications |
|--------|------------------------|--------------|
| **Hypoglycemia risk** | Older adults are at increased risk of glycemic variability, hypoglycemia, and hypoglycemia unawareness, leading to greater risk for negative outcomes from hypoglycemic events [1–10, 60–62] | Prioritize hypoglycemia prevention Avoid intermediate-acting insulin formulations for basal control Consider expanding therapeutic options to long-acting injectable agents with demonstrated efficacy and low risk of hypoglycemia Administer basal insulin in the morning when concerns for hypoglycemia are highest Long-acting GLP-1 RA and basal insulin formulations are effective and safe | Fewer injections per day may also counter patient-related barriers, reducing issues from those related to the injection itself (such as discomfort) to those countering lack of function and dependence on others to administer the medication |
| **Patient-related barriers** | Individual (fear of pain and injections, concerns regarding side effects, social stigma) [59] Provider (knowledge and skills, clinical inertia, language) System (time, education resources) | Patient and provider education is paramount Informed consent discussion and common agreement to the plan of care (with patient and caregiver if appropriate) is crucial Pertinent monitoring, follow-up, and support, with a chronic disease self-management approach, are important to secure ongoing adherence, address questions and issues, and adjust glycemic targets and plan as needed Ongoing education and updates on newer strategies are recommended; this may include training in motivational interviewing for the main provider and the clinic staff engaged in the diabetes management of the older patient Education and training on the use of newer injectable agents (for providers and their staff) may enhance treatment strategies and expand the options to be presented to older patients with T2D |
| **Clinical inertia** | Barriers to treatment intensification [63] It is highly prevalent [64] Poor glycemic control can lead to complications [65, 66] Older age itself can be an added risk factor for clinical inertia [67] | |

*GLP-1 RA* glucagon-like peptide-1 receptor agonist, *T2D* type 2 diabetes
4.2 Reducing Patient-Related Barriers

Providers must identify the reasons for patients’ ambivalence towards treatment, weighing views regarding advantages and disadvantages, and set clear goals and strategies to overcome the treatment barriers. Trained coaches can deliver an intervention at the point of care or via telephone.

All patients should be taught that as diabetes progresses, they may require basal insulin and, in the future, they might also need bolus-insulin dosing to better manage mealtime hyperglycemia exposure [9].

Improved patient and caregiver education are crucial for ensuring the successful transition to injectable therapies. Attention should be paid to dosage and self-management [60]. Interventions should include hands-on demonstrations of insulin injection, together with increased support from healthcare providers, using trained staff to remove time constraints, and regular follow-up via clinic visits or telephone calls.

In parallel, cognitive behavior strategies, such as motivational interviewing, may also help patients to improve self-efficacy and adherence to chronic medications [76]. Providers and staff may be trained to improve delivery of interventions to patients [60].

Previous reports indicate that pen-delivery devices are effective, improve convenience and flexibility, and may improve patients’ confidence [77]. While the reduction of social stigma is still relevant for older adults, a greater advantage of pen delivery over the vial-and-syringe method is the greater ease of use for those with decreased physical dexterity. A database study in older adults with T2D using insulin pens compared with vial-and-syringe administration reported improved adherence and persistence with insulin therapy in the pen group [78].

Remarkably, despite the higher drug costs for pen devices, a claims database analysis did not find increased total all-cause or diabetes-related healthcare costs when compared with vial and syringe and, additionally, reported lowering of hypoglycemia rates [79]. The latter could decrease costs related to emergency-room visits and hospitalizations.

Despite the lack of studies focused on older adults in specific settings, we have previously suggested that newer, once-weekly injectable GLP-1 RAs may be used instead of long-acting insulin in older patients with impaired mobility, limited social support, or a combination of these [1]. DURATION-3 was an RCT that compared once-weekly exenatide with daily Gla-100 in 456 patients with an average age of 58 years [80]. The researchers found a modest advantage (0.16% greater reduction in A1C) in favor of the long-acting exenatide, yet we note their success in implementing a non-insulin injectable agent required only once-weekly administration in addition to oral agents to accomplish glycemic targets. The 3-year follow-up of these patients indicated sustained efficacy, which could enhance adherence and reduce barriers by reducing the number of injections and services required [81].

When treatment requires intensification, injectable therapies must be balanced with the treatment burden and the consequences of a delayed or missed insulin dose [11]. A ‘basal–bolus’ regimen (i.e., basal insulin and short-acting insulin before meals) adds up to at least four injections and four fingerstick fasting serum glucose measurements per day. This can be burdensome for both independent older adults as well as for those who rely on caregivers for their management. The availability of newer, improved (less thumb force and time needed to inject insulin) pen needles can mitigate the discomfort due to injections and is associated with improved patient satisfaction [82]. Thus, newer fixed-ratio combination agents, IDegLira or iGlarLixi, could reduce the number of injections while offering basal glycemic control and postprandial glucose control from the GLP-1 RA. In parallel, educational interventions and the use of telehealth technology can help overcome barriers to adherence and resistance to therapy. Strategies may include health coaching interventions, with or without the use of mobile-phone monitoring as support.

4.3 Reducing Clinical Inertia

Individualized targets and understanding of the patient’s preferences and prognosis can provide settings where the next step of management involves de-intensification of therapies on the one hand, but an actual need to intensify therapy in those healthy older adults who may live long enough to suffer from complications due to poorly controlled diabetes, and may help create awareness regarding the heterogeneous clinical presentation of diabetes in older adults, even within the same age brackets [1, 3, 83]. Although referral to an endocrinologist was found to be associated with a reduced risk of clinical inertia [66], general practitioners may intensify therapies accordingly. Continued medical education regarding emerging treatment strategies and different formulations may enhance the informed discussion with patients and reduce clinical inertia. Providers will benefit from understanding the potential advantages of newer agents, including lower risk of nocturnal hypoglycemia, fewer injections (daily, weekly), feasible titration of dosages, as well as the potential disadvantages, especially costs [84, 85], access, and barriers to injectables in general, as described above. In addition, using telehealth for remote monitoring is another potential strategy to improve management and reduce clinical inertia, but we did not find clinical trials focused on older adults with complex clinical scenarios.

Discussing diabetes care is a complex endeavor. It should also be considered that due to the natural history of the disease, as well as its chronicity and progression, older adults
tend to require a greater number of medications (including exogenous insulin) to achieve glycemic control. We must acknowledge the potential risk of negative consequences of overtreatment, which may include polypharmacy, hypoglycemia risk, treatment burden, side effects, and costs. All these factors should be considered and balanced when selecting antihyperglycemic agents. Finally, based on our clinical practice, when we discuss the use of injectable agents, especially insulin-based products, many patients raise the concern about the long-term effects. They are worried about the common understanding that once a patient starts insulin therapy, there is no turning back. There are multiple approaches to address this situation, taking into account that diabetes is chronic and progressive.

1. Proper counseling throughout the course of the disease, even while well controlled, education and counseling on the ‘worst-case scenarios’ (e.g., standard practice when A1C is greater than 11%) and that diabetes progression is expected, so that we do not know what the pharmacologic needs will be in 3, 5, 10, 20, or more years.

2. Proper education and monitoring, even when patients are not ready to start an injectable agent, which they might well need. Always leave the opportunity open to resume the conversation, understanding and addressing what are the specific concerns they may have, and build a plan of action. This often involves giving them the opportunity to try, on their own, things they want to do, for example, “give me 3 months to improve my lifestyle”; this can be plausible if, during the visit, we detect there is actual room for improvement.

3. Let patients know that starting an injectable, such as insulin, does not imply the treatment is set in stone. Notably, patients can improve weight and lifestyle, and perhaps adherence to other medications, and while this may not occur for all, it is possible that some can make significant improvements to the point that active downtitration of injectables is required to reduce hypoglycemia risk. We have observed patients in whom the reduction of insulin dosages reaches the point where we do not observe any impact (e.g., when short-acting insulin has to be titrated below 3 units in a patient with T2D, who previously had obesity stage 2, and then reduced their weight to the overweight category).

5 Conclusions

The evaluation and management of T2D in older adults is complex and challenging as the disease can impact all four geriatric domains. Working with patients and involving their caregivers is crucial to ensure proper clinical care, especially in those with functional or cognitive decline who require formal or informal assistance. There is a well-established concern regarding hypoglycemia, but hyperglycemia/uncontrolled diabetes can also lead to multiple complications in this age group. Clinicians should implement strategies to overcome barriers and clinical inertia, to ensure older patients reach their individualized glycemic targets. For some patients, injectable agents may offer advantages that can overcome barriers, including lower risk of hypoglycemia, effective glycemic control, and available weekly formulations. Unfortunately, there is a paucity of dissemination strategies to incorporate injectable agents in this age group, which makes this practice difficult.

We strongly recommend early patient education on the chronicity and progressive characteristics of this disease, as well as preparing patients for their potential future need for injectable therapies. While the newest combined agents (long-acting insulin with GLP-1 RA in one injection) have not been extensively investigated in older adults in the community or in long-term care, they could potentially be implemented at the primary care level and in long-term care, instead of basal–bolus regimens, as they may offer potential advantages, including greater adherence and less treatment burden. We recommend long-term prospective studies to monitor patient-centered (quality of life, prevention of complications, hypoglycemia, emergency room visits, and hospitalizations) and system-centered (cost-effectiveness, reduced need for skilled nursing services and formal support) outcomes, comparing different injectable agents and strategies in older adults living in the community and long-term care settings. In the meantime, the available evidence shows that implementing injectable therapies in older adults can be feasible, effective, and potentially safer than more traditional agents. Further education and engagement of general practitioners and their staff are paramount.

Costs are and should always be considered when selecting agents. However, clinical inertia should not be due to cost issues as many options are available.

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Compliance with Ethical Standards

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