



Consensus guidance for monitoring individuals with islet autoantibody-positive pre-stage 3 type 1 diabetes

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Abstract

Given the proven benefits of screening to reduce diabetic ketoacidosis (DKA) likelihood at the time of stage 3 type 1 diabetes diagnosis, and emerging availability of therapy to delay disease progression, type 1 diabetes screening programmes are being increasingly emphasised. Once broadly implemented, screening initiatives will identify significant numbers of islet autoantibody-positive (IAB⁺) children and adults who are at risk of (confirmed single IAB⁺) or living with (multiple IAB⁺) early-stage (stage 1 and stage 2) type 1 diabetes. These individuals will need monitoring for disease progression; much of

This consensus report was endorsed by the European Society for the Study of Diabetes (EASD), American Diabetes Association (ADA), American Association of Clinical Endocrinology (AACE), American College of Diabetology (ACD), Association of Diabetes Care & Education Specialists (ADCES), Australian Diabetes Society (ADS), the International Society for Pediatric and Adolescent Diabetes (ISPAD), Advanced Technologies & Treatments for Diabetes (ATTD), DiaUnion, the Endocrine Society and JDRF International.

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A consensus report of a particular topic contains a comprehensive examination and is authored by an expert panel and represents the panel's collective analysis, evaluation and opinion. Consensus reports are reviewed for EASD by its Committee for Clinical Affairs (CCA). The CCA may, at their discretion, enlist further external expert reviewers.

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this care will happen in non-specialised settings. To inform this monitoring, JDRF in conjunction with international experts and societies developed consensus guidance. Broad advice from this guidance includes the following: (1) partnerships should be fostered between endocrinologists and primary-care providers to care for people who are IAb⁺; (2) when people who are IAb⁺ are initially identified there is a need for confirmation using a second sample; (3) single IAb⁺ individuals are at lower risk of progression than multiple IAb⁺ individuals; (4) individuals with early-stage type 1 diabetes should have periodic medical monitoring, including regular assessments of glucose levels, regular education about symptoms of diabetes and DKA, and psychosocial support; (5) interested people with stage 2 type 1 diabetes should be offered trial participation or approved therapies; and (6) all health professionals involved in monitoring and care of individuals with type 1 diabetes have a responsibility to provide education. The guidance also emphasises significant unmet needs for further research on early-stage type 1 diabetes to increase the rigour of future recommendations and inform clinical care.

Keywords Autoantibodies · Glucose monitoring · Prevention · Type 1 diabetes

Abbreviations

ADCES	Association of Diabetes Care & Education Specialists
CGM	Continuous glucose monitoring
DCES	Diabetes care and education specialists
DKA	Diabetic ketoacidosis
DSMES	Diabetes self-management education and support
HCP	Healthcare professional
IAb ⁻	Islet autoantibody negative
IAb ⁺	Islet autoantibody positive
IASP	Islet Autoantibody Standardisation Program
ISPAD	International Society for Pediatric and Adolescent Diabetes
SMBG	Self-monitored blood glucose
TEDDY	The Environmental Determinants of Diabetes in the Young

Overview

Currently, screening of individuals for islet autoantibodies is undertaken as part of programmes to detect children, adolescents and adults who are at higher risk of developing type 1 diabetes due to having a first-degree relative with type 1 diabetes or having a known high-risk HLA genotype. Periodic monitoring of people who have screened positive for one or more autoantibodies (islet autoantibody-positive [IAb⁺] individuals) is largely, but not always, conducted within these cohort studies. However, up to 90% of people who develop type 1 diabetes are not part of at-risk groups. Thus, screening programmes within the general population are being initiated and guidance for monitoring in non-specialist settings is urgently needed. The guidance provided here was developed by a series of expert working groups, convened as part of a JDRF initiative to document the aims, scope and purpose of monitoring for children, adolescents and adults with islet autoantibody positivity, along with recommended

frequencies of monitoring and actions for healthcare professionals (HCPs) when risk of progression towards symptomatic type 1 diabetes is high. This includes expert clinical advice for educational and psychosocial support for IAb⁺ individuals, including for their families and caregivers. The expert clinical advice for adults reflects available data, yet it is important to note that there are very limited data in adults aged 45 years and older who are IAb⁺. It is also important to note that this consensus document does not encompass screening for islet autoantibodies, and only provides expert clinical advice for monitoring of individuals who have screened positive for at least one islet autoantibody.

Introduction and rationale

The presence of islet autoantibodies for a presymptomatic period of variable duration in first-degree relatives of individuals with type 1 diabetes has been known for more than 40 years [1], with recommendations for islet autoantibody screening appearing soon after [2]. Decades of subsequent research and monitoring of individuals with islet autoantibody positivity has led to the paradigm shift that type 1 diabetes is a continuum of stages, from genetic risk through to autoimmunity and then metabolic disease. This has been accompanied by the evolution of descriptive terminology that reflects these stages (Table 1). Similarly, treatment options have moved on from monitoring and managing metabolic disease to include options for modulating the autoimmune response [3, 4].

Screening programmes have developed to the point that large numbers of children and adults at risk of and with early-stage type 1 diabetes have been intensively followed in longitudinal cohort studies [5–15] centred on understanding the natural history of progression to symptomatic type 1 diabetes (see Table 2 for a list of studies

Table 1 Staging criteria for autoantibody-positive individuals in pre-stage 1 and stage 1–3 type 1 diabetes [16–18]

Stage of T1D	Islet autoantibody status	Glycaemic status	Symptoms	Insulin required
At-risk (pre-stage 1 T1D)	Single autoantibody or transient single autoantibody	<ul style="list-style-type: none"> • Normoglycaemia • FPG <5.6 mmol/l (<100 mg/dl) • 120 min OGTT <7.8 mmol/l (<140 mg/dl) • HbA_{1c} <39 mmol/mol (<5.7%) 	No symptoms	Not required
Stage 1 T1D (also referred to as early-stage T1D or presymptomatic T1D)	≥2 autoantibodies	<ul style="list-style-type: none"> • Normoglycaemia • FPG <5.6 mmol/l (<100 mg/dl) • 120 min OGTT <7.8 mmol/l (<140 mg/dl) • HbA_{1c} <39 mmol/mol (<5.7%) 	No symptoms	Not required
Stage 2 T1D (also referred to as early-stage T1D or presymptomatic T1D)	≥2 autoantibodies ^a	<p>Glucose intolerance or dysglycaemia not meeting diagnostic criteria for stage 3 T1D, with at least two of the following, or meeting the same single criteria at two time points within 12 months:</p> <ul style="list-style-type: none"> • FPG 5.6–6.9 mmol/l (100–125 mg/dl) • 120 min OGTT 7.8–11.0 mmol/l (140–199 mg/dl) • OGTT values ≥11.1 mmol/l (≥200 mg/dl) at 30, 60 and 90 min • HbA_{1c} 39–47 mmol/mol (5.7–6.4%) or longitudinal ≥10% increase in HbA_{1c} [66, 67] from the first measurement with stage 2 T1D • CGM values >7.8 mmol/l (>140 mg/dl) for 10% of time over 10 days' continuous wear [73]^b, and confirmed by at least one other non-CGM glucose measurement test listed 	No symptoms	Not required
Stage 3 T1D	≥1 autoantibody	<p>Persistent hyperglycaemia with or without symptoms, as measured and confirmed by one or more of the following:</p> <ul style="list-style-type: none"> • one random venous glucose ≥11.1 mmol/l (≥200 mg/dl) with overt symptoms • 120 min OGTT ≥11.1 mmol/l (≥200 mg/dl) and/or • two random venous glucose ≥11.1 mmol/l (≥200 mg/dl) and/or • FPG ≥7.0 mmol/l (≥126 mg/dl) and/or • laboratory-tested HbA_{1c} ≥48 mmol/mol (≥6.5%) • CGM values >7.8 mmol/l (>140 mg/dl) for 20% of time over 10 days' continuous wear [73]^b and confirmed by at least one other non-CGM glucose measurement test listed 	<p>May include^c:</p> <ul style="list-style-type: none"> • Polyuria • Polydipsia • Weight loss • Fatigue • DKA 	+/- Insulin, based on glycaemic status

^aSome people with confirmed persistent prior multiple autoantibody positivity may revert to single autoantibody status or negative status [95]

^bCGM is ideally blinded and must be applied and interpreted by a trained HCP. Note, use of CGM-derived criterion did not achieve consensus within the consensus panel and CGM metrics are not part of current ADA or ISPAD guidelines on staging criteria in type 1 diabetes [16, 155]

^cStage 3 might not include symptoms

FPG, fasting plasma glucose; T1D, type 1 diabetes

available for participation). Of note, many entry criteria for individuals with presymptomatic type 1 diabetes into these studies require a family history of type 1 diabetes or HLA genetic risk, and most are focused on paediatric populations. Based on the outcomes of these and other studies, stages of presymptomatic and symptomatic type 1 diabetes are now clinically defined (Table 1) to a degree of clinical consensus [16–18], although regulatory agencies and research studies may differ in definitions.

Using these classifications, individuals can be monitored, diagnosed with diabetes and even, at times, started on insulin replacement therapy early in the disease course, based on meeting American Diabetes Association (ADA) [18], International Society for Pediatric and Adolescent Diabetes (ISPAD) [16] or American Association of Clinical Endocrinology (AACE) [19] diagnostic criteria. To date, the ISPAD guidelines have provided metabolic and autoantibody monitoring recommendations for children

Table 2 Established population-based screening and monitoring studies in early-stage type 1 diabetes

Acronym	Study name/description
ASK	Autoimmunity Screening for Kids programme [7]
BABYDIAB	Part of the international Type 1 Data Intelligence (T1DI) project [156]
DAISY	Diabetes Autoimmunity Study in the Young [6]
DIPP	Type 1 Diabetes Prediction and Prevention Study based in Finland [11]
DPT-1	Diabetes Prevention Trial–Type 1 [12]
ENDIT	European Nicotinamide Diabetes Intervention Trial [13]
Fr1da	Population-based healthcare research study based in Bavaria, Germany [9]
INNODIA	Global partnership between academic institutions, commercial partners and patient organisations [14]
PLEDGE	Population Level Estimation of T1D Risk Genes in Children [155]
TEDDY	The Environmental Determinants of Diabetes in the Young study [5]
Type 1 Diabetes TrialNet	International research network centred on delaying or preventing T1D [10]
Type1Screen	Australian screening and monitoring programme open to relatives of individuals with type 1 diabetes and IAb ⁺ people identified through other screening pathways (ANZCTR registration no. ACTRN12620000510943)

Note that major research networks are included in the table, but this is not an exhaustive list
ANZCTR, Australian New Zealand Clinical Trials Registry

with presymptomatic type 1 diabetes [16], but do not make specific recommendations for education or psychosocial support in IAb⁺ individuals, monitoring of single IAb⁺ individuals or when to start insulin. The Fr1da study has suggested and introduced specific recommendations for children [20]. A separate set of recommendations based on a Delphi-survey of expert opinion has provided guidance on metabolic and autoantibody monitoring, with recommendations for education and

psychosocial support, but does not specifically address adults with early-stage type 1 diabetes [21]. Consequently, to date there is no available guidance on monitoring in adults or in individuals with single islet autoantibody positivity, or on when insulin therapy is indicated.

Consensus on evidence-based expert clinical advice for monitoring is an important unmet need since a positive test for islet autoantibodies (Table 3) is a condition for access to disease-modifying therapies, such as teplizumab [22].

Table 3 Autoantibodies against islet autoantigens detected in stage 1–3 type 1 diabetes

Autoantibody	Islet specificity	Typical characteristics
IAA	Insulin	<ul style="list-style-type: none"> • Common as a first detected autoantibody in young children [157, 158] • Appearance is more common in younger children [159] • Frequency of appearance declines with age • Not informative for individuals treated with insulin, who often develop antibodies in response to injected insulin
GADA	GAD	<ul style="list-style-type: none"> • Common as a first detected autoantibody in childhood, up until age 15 years [157, 158, 160] • Adult-onset cases most often present with GADA [161] • Is associated with slower progression to T1D [162] and is often found as a single positive islet autoantibody, especially in adults
IA-2A (also known as ICA512)	Tyrosine phosphatase islet antigen-2	Presence is associated with more-advanced islet autoimmunity and faster progression to stage 3 T1D [55, 163]
ZnT8A	Zinc transporter type 8, a trans-membrane protein in the beta cell granule	Presence can improve risk stratification in individuals with single GADA ⁺ , IAA ⁺ or IA-2A ⁺ status [164]
ICA	Multiple antigens, undefined	Detected by indirect immunofluorescence on islet cell tissue. While not frequently measured other than in research studies, it does add to risk determination in the presence of other biochemical autoantibodies

IA-2A, insulinoma antigen-2 autoantibody; ICA512, islet cell autoantigen 512; T1D, type 1 diabetes

In addition, islet autoantibody screening is anticipated to become more common [7, 23–25], highlighting the need for clear monitoring advice.

Screening efforts are identifying an ever-growing number of IAb⁺ people who warrant education and ongoing monitoring for progression towards clinical diabetes. Evidence shows that such monitoring in research studies can significantly reduce the incidence of diabetic ketoacidosis (DKA) at diagnosis [24, 26–33], occurring in up to 70% of unmonitored individuals, which is greatly lowered for individuals participating in follow-up studies [26, 34–39]. The impact of monitoring in general clinical practice on DKA rates is not known. DKA is a life-threatening condition that requires hospital admission, with significant associated costs for critical care [40–42]. Additionally, in a number of studies, DKA at presentation of type 1 diabetes in youths has been associated with higher HbA_{1c} that was sustained for up to 11 years after diagnosis [43–45]. Other studies have, however, not found such an association between DKA at presentation of type 1 diabetes and higher long-term glycaemic levels [46]. The lack of DKA at onset of type 1 diabetes is also predictive of fewer severe hypoglycaemic events 10 years after diagnosis [47]. In this context, the overall goals of monitoring are described in Table 4.

Monitoring of people with islet autoantibody positivity outside of research settings will require expert clinical advice that is clear and actionable by HCPs who have limited expertise in diabetes. As indicated, current insights into monitoring progression to clinical type 1 diabetes are largely derived from research studies of individuals known to be at risk of type 1 diabetes, and general population data are less extensive. With this caveat, knowledge on best practices is particularly important for primary-care and secondary-care physicians who may not frequently see people known to be at risk of type 1 diabetes, and yet who will be tasked with the initial aspects of monitoring following a positive autoantibody screen. Other people who may assist with care of these individuals will include nurse practitioners, physician assistants, diabetes care and education specialists (DCES), psychologists and other mental and behavioural health professionals, all of whom have a role in supporting IAb⁺ individuals and their families within the

monitoring environment. Clear expert clinical advice for monitoring by these groups of HCPs increases the likelihood that individuals at risk for or in early stages of type 1 diabetes, and their families, can receive accurate and actionable education about presymptomatic type 1 diabetes and their individual status.

The requirement for monitoring

Islet autoantibodies against four major pancreatic autoantigens are currently clinically available; these consist of IAA, GADA, insulinoma antigen-2 autoantibody (IA-2A; also called islet cell autoantigen 512 [ICA512]) and ZnT8A [48]. These are often considered ‘biochemical autoantibodies’ and are the screening targets recommended by the most-recent ADA Standards of Care [25]. A further islet autoantibody assay, for ICA, using indirect immunofluorescence on pancreatic tissue, has been used for screening purposes, but it is less available outside of research studies and the antigenic targets are not fully known. Considerable evidence in multiple populations supports the concept that the number and type of biochemical autoantibodies can be used to predict risk for progression to clinical disease (stage 3 type 1 diabetes; see Table 1). These autoantibodies and their characteristics are described in Table 3. However, it must be noted that these attributes are derived from observations made in known IAb⁺ populations in the research environment. Further data from studies in IAb⁺ groups in the general population are needed.

Confirmation of IAb⁺ status is important to identify the persistence of the underlying autoimmune response and the validity of the target antigen, although the accuracy of autoantibody tests can vary between laboratories and between target antigens. Therefore, the first positive test should be confirmed with a second test within 3 months [49] and, where possible, in a laboratory that meets the performance standards set by the Islet Autoantibody Standardization Program (IASP) [50]. Persistent IAb⁺ status on two or more different samples is needed, using sensitive and specific assays with high predictive value for disease progression [51]. Several research programmes have tested for islet autoantibody status using capillary sampling to obtain

Table 4 Purpose of monitoring in IAb⁺ children, adolescents and adults

1. Primary purpose is to prevent DKA and to minimise the risk of requiring emergency care or hospital admission
2. Identification for and monitoring of therapeutic intervention(s) to delay stage 3 T1D onset (where available) and prolong beta cell function
3. To provide advice for the start of insulin in stage 3 T1D, when glucose is sufficiently elevated and before symptoms develop, to optimise HbA_{1c} and avoid the consequences of hyperglycaemia on long-term glycaemic outcomes
4. To avoid misdiagnosis of T2D and delayed commencement of insulin therapy
5. Referral for participation in research studies

T1D, type 1 diabetes; T2D, type 2 diabetes

Table 5 Attributes of current monitoring methods

Method	Pros	Cons	Metrics obtained
Reference OGTT ^a	<ul style="list-style-type: none"> • Gold standard in research settings • Used to stage disease and predict progression 	<ul style="list-style-type: none"> • Requires glucose load and 2–5× blood draws over 2 h 	<ul style="list-style-type: none"> • Glycaemic staging • Risk scores for progression (DPTRS, DPTRS60, Index60, M60, M120, PLS) [94, 165–169]
Standard OGTT ^b	<ul style="list-style-type: none"> • Similar to test for GDM: OGTT with 2× blood draws (compared with 3× draws in GDM test), performed routinely in clinical care • One-off sample • Low cost • Highly specific for clinical diagnosis of stage 3 T1D • Can use capillary sample • Longitudinal HbA_{1c} may be as informative as OGTT [66] 	<ul style="list-style-type: none"> • Requires 2× blood draws: fasting and at 2 h 	<ul style="list-style-type: none"> • 120 min OGTT-derived glucose • M120
Random glucose	<ul style="list-style-type: none"> • One-off sample • Low cost 	<ul style="list-style-type: none"> • Requires a blood draw or fingerstick test • Less sensitive than 120 min OGTT 	<ul style="list-style-type: none"> • Similar to 120 min OGTT-derived glucose [96] if obtained 2 h postprandially
Standard HbA _{1c} test	<ul style="list-style-type: none"> • Highly specific for clinical diagnosis of stage 3 T1D • Can use capillary sample • Longitudinal HbA_{1c} may be as informative as OGTT [66] 	<ul style="list-style-type: none"> • Indicates 3 month mean glucose. Often normal in asymptomatic or recent-onset stage 3 T1D • May be affected by age, non-diabetes disease states (e.g. renal, haematological syndromes) • Not suitable in the home setting 	<ul style="list-style-type: none"> • Risk of progression to ‘clinical disease’: HbA_{1c} >39 mmol/mol (>5.7%) [170] • 10% rise from baseline (at first positive islet autoantibody) over 3–12 months [66, 67], suggests dysglycaemia and progression to stage 2 T1D • Consider use of CGM if 10% rise in HbA_{1c} is confirmed, or higher frequency of SMBG, to monitor risk for progression
CGM ^c	<ul style="list-style-type: none"> • Can be used at home • Can be blinded for physician review only in some regions • Optimal duration of CGM wear is validated in adults and children >2 years of age with diagnosed T1D, at all glycaemic levels [171] 	<ul style="list-style-type: none"> • Risk of anxiety for unblinded user seeing CGM fluctuations and experiencing alarms • Requires appropriate education on use and interpretation • Many primary-care HCPs are unfamiliar with interpretation • Cost and access issues • Duration of wear not validated in early-stage T1D 	<ul style="list-style-type: none"> • Sensitive in detecting individuals with asymptomatic stage 3 T1D and dysglycaemia in stage 2 T1D [73] • Risk of progression to ‘clinical disease’, i.e. 10% of time with glucose >7.8 mmol/l (>140 mg/dl) has been associated with an 80% risk of progression to T1D within 12 months [72] • ≥5% time with glucose ≥7.8 mmol/l (≥140 mg/dl) has been associated with a 40% risk of progression to T1D within 2 years [71] • Other PPV metrics not tested
SMBG	<ul style="list-style-type: none"> • Simple to use at home • Comparatively low cost 	<ul style="list-style-type: none"> • Uncomfortable for users, can affect accuracy and use • Optimal timing and frequency have not been determined 	<ul style="list-style-type: none"> • Immediate capillary blood glucose test result • 2 h postprandial measure likely of most value
C-peptide	<ul style="list-style-type: none"> • Validated measure of beta cell function • Stimulated C-peptide in research settings is valuable to assess insulin production and distinguish between T1D (or stages of T1D) and T2D 	<ul style="list-style-type: none"> • Can be falsely low in hypoglycaemia <3.9 mmol/l (<70 mg/dl), in severe hyperglycaemia/DKA or after fasting, so concomitant serum glucose should be checked for interpretation • Wide range of values at clinical diagnosis, including >0.2 nmol/l, and persistent, but low, levels of secretion can be seen long after diagnosis • Presence of C-peptide does not exclude T1D and on its own is not useful for staging or diagnosis of T1D 	<ul style="list-style-type: none"> • A stimulated postprandial C-peptide value ≤0.2 nmol/l with IAb⁺ status can assist with appropriately classifying diabetes type
Repeat antibody testing	<ul style="list-style-type: none"> • Confirms initial IAb⁺ test result and progression to multiple IAb⁺ status 	<ul style="list-style-type: none"> • None 	<ul style="list-style-type: none"> • Autoantibody type and single IAb⁺ or multiple IAb⁺ status
Education	<ul style="list-style-type: none"> • Provides awareness of diabetes symptoms and signs 	<ul style="list-style-type: none"> • None 	<ul style="list-style-type: none"> • Person-reported outcomes for possible progression to stage 3 T1D

^aUsed in research settings for staging progression of impaired glucose tolerance as C-peptide provides important predictive value

^bUsed in clinical practice to detect impaired glucose tolerance in prediabetes and gestational diabetes mellitus

^cUse of CGM-derived criterion did not achieve consensus within the consensus panel, with further evidence required to confirm findings to date

DPTRS, Diabetes Prevention Trial-Type 1 risk score; GDM, gestational diabetes mellitus; M60, 60 min test result; M120, 120 min test result; PLS, partial least squares; PPV, positive predictive value; T1D, type 1 diabetes; T2D, type 2 diabetes

serum or dried blood spots for assessment; however, venous samples are preferred (due to reduced interference from haemolysis) and should be used as confirmation whenever capillary testing has been performed initially.

Predicting when an individual with type 1 diabetes-related autoantibodies may progress to stage 3 type 1 diabetes is difficult. However, in children and adolescents, persistent multiple IAb⁺ status confirms early-stage (stage 1 or stage 2) type 1 diabetes with higher rate of progression to stage 3 type 1 diabetes compared with single IAb⁺ status [52]. For the same reasons as discussed for single IAb⁺ status, confirmation of multiple IAb⁺ status is important, as it indicates early-stage type 1 diabetes, and should adhere to the ‘rule of twos’, i.e. the presence of two different autoantibodies, confirmed in two tests from two separate samples [51–54]. Subsequent loss of individual antibodies is not associated with a slower rate of progression. The type of positive autoantibody (Table 3) is also of importance, since, as children age, relative risks for progression with each antibody type will change [55, 56], with some evidence that this is also true for adults [55, 57]. Consideration of these data, along with autoantibody titres, may aid risk stratification [58]. Although fewer data are available in adults, Type 1 Diabetes TrialNet cohort data indicate that the rate of progression to type 1 diabetes in IAb⁺ adults is slower than in children [59].

Misdiagnosis of type 1 diabetes as type 2 diabetes in adolescents and adults can lead to DKA [60], as this misdiagnosis means that these individuals are often not started on insulin [61]. Latent autoimmune diabetes of adults (LADA) can also be misdiagnosed as type 2 diabetes [62], with a risk of delayed insulin initiation. These observations emphasise the value of autoantibody testing for newly-diagnosed adults with diabetes, particularly when they have features of type 1 diabetes (e.g. younger age, non-obese, sudden weight loss, mild acidosis, DKA, hyperglycaemia > 16.7 mmol/l [>300 mg/dl]) [63], for making an accurate diagnosis and starting appropriate treatment. It is, however, important to recognise that some individuals with new-onset type 1 diabetes have a phenotype that does not differ substantially from people with type 2 diabetes, particularly given the increased prevalence of obesity [60, 64]. Misdiagnosis of MODY is also reported [65], suggesting that islet autoantibody screening can be valuable at presentation of all forms of diabetes.

An important outcome of monitoring individuals with islet autoantibody positivity is to inform the decision to initiate insulin therapy, and this is an area of evolving practice. In some centres, individuals with hyperglycaemia (see Table 5) but with HbA_{1c} <48 mmol/mol (<6.5%) might not be started on insulin without the presence of symptoms. Sequential HbA_{1c} monitoring has been productive in this context in paediatric studies on individuals

with islet autoantibody positivity, since an absolute $\geq 10\%$ increase from baseline, even if the HbA_{1c} test reading stays below 48 mmol/mol (6.5%), is predictive of disease progression [66, 67] within a median of 1 year. Risk of progression within 2 years following a confirmed $\geq 10\%$ increase in HbA_{1c} is lower for older individuals. This aspect of stage 3 type 1 diabetes (i.e. when to start insulin once hyperglycaemia is confirmed) requires further evidence to support clinical practice, to better understand the metabolic and mental-health outcomes.

What should be monitored? It is acknowledged that the practice of monitoring of individuals with islet autoantibody positivity must accommodate different settings with diverse healthcare resources. In this context, there are multiple available tools for monitoring, including self-monitored blood glucose (SMBG), periodic continuous glucose monitoring (CGM), a standard OGTT, random venous glucose, HbA_{1c} and repeat islet autoantibody monitoring. In this context, serial stimulated C-peptide measurement during an OGTT can be used to assess deterioration of beta cell function and to predict risk development of type 1 diabetes [68]. Since individuals who present with clinical type 1 diabetes (stage 3) often have significant residual beta cell function [69], they may benefit from therapies that can optimise prolongation of insulin secretion [70].

The pros and cons of each monitoring method are documented in Table 5. Identification of an increase in sequential HbA_{1c} values from a baseline reading can be as informative as 2 h OGTT values in predicting risk of stage 3 type 1 diabetes in youth with genetic risk and type 1 diabetes-associated autoantibodies [66, 67]. Ongoing research continues to evaluate the role of CGM (including professional CGM, which is blinded to the user) in aiding in the identification of individuals, including those with a normal OGTT, who are likely to rapidly progress to stage 3 type 1 diabetes [71–73]. To date, use of CGM metrics in individuals who have multiple IAb⁺ status has been shown to be predictive of progression to type 1 diabetes, but CGM measures are not yet as sensitive as OGTT testing [74].

Where should monitoring take place? In practice, monitoring should be carried out wherever the skills and resources exist to perform the appropriate tests (Table 5). However, since many people will be monitored in primary care, there is a need to consider different intensities of monitoring consistent with resources available. The capabilities of primary-care HCPs and other care providers should be applied to monitoring of early-stage type 1 diabetes without the need to refer to an expert practitioner, until clinically appropriate. In primary care, this may help specify basic education about symptoms and glycaemic signposts.

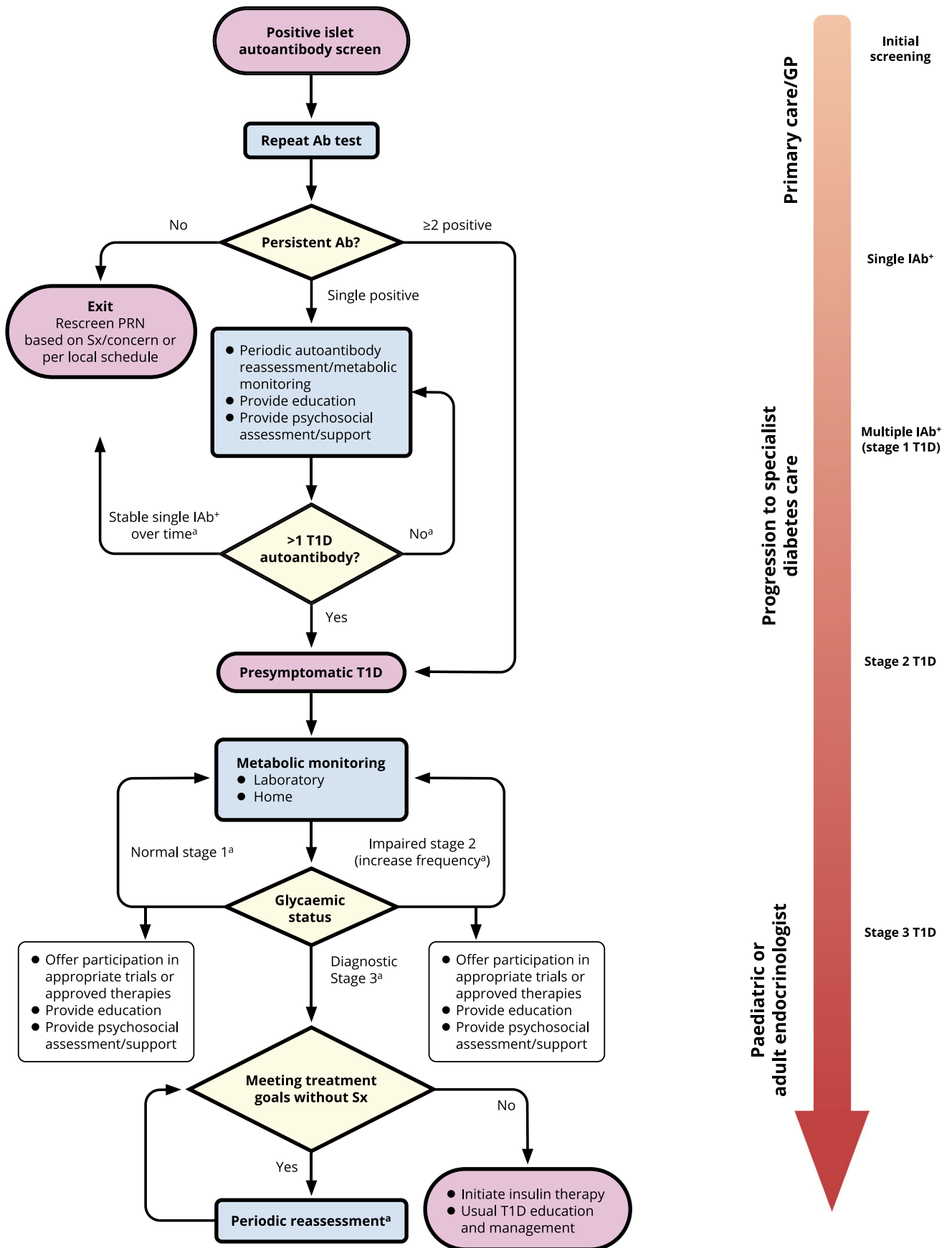


Fig. 1 Algorithm for monitoring of people screened positive for one or more islet autoantibodies. ^aMonitoring frequency and methodology depends on age, length of time since first detection of islet autoantibody, number of islet autoantibodies detected and presence of symptoms of type 1 diabetes (see Tables 1, 3, 4 and 5). Ab, antibody; GP, general practitioner; PRN, pro re nata (as needed); Sx, symptoms; T1D, type 1 diabetes. This figure is available as part of a [downloadable slideset](#)

It is understood that, compared with stage 1, monitoring in stage 2 type 1 diabetes may require more-expert practitioners.

Objectives and methodology

The aim of this international consensus report is to formulate expert clinical advice, based on current evidence and expert opinion, that specifies the required monitoring and management approach for people who have been identified as having IAb⁺ status and pre-stage 3 type 1 diabetes, and can be used in daily clinical practice. Overall, these key principles should encompass: (1) who should be monitored; (2) which endpoints to monitor; (3) the frequency and duration of monitoring; (4) initiation of insulin during stage 3 type 1 diabetes; and (5) how to provide psychosocial and educational support for affected individuals and families.

We acknowledge that monitoring of IAb⁺ individuals will occur in diverse settings, with variable resources to support effective monitoring of IAb⁺ individuals. Thus, a guiding principle of this consensus report is to provide advice that is straightforward and actionable within the landscape of available clinical skills and resources, wherever the monitoring will take place. The audience for this consensus document, therefore, includes: (1) primary-care providers; (2) endocrinologists and diabetologists; (3) DCES; (4) mental and behavioural health professionals; and (5) individuals at risk for or in early stages of type 1 diabetes and their families.

Methodology

The consensus process was initiated by the JDRF with a conference held on 21 February 2023 at the 16th International Conference on Advanced Technologies & Treatments for Diabetes (ATTD) in Berlin, Germany, with in-person or virtual attendance. MP served as Chair of

the project and LAD served as Vice Chair. A mission statement was created and the attendees were invited by email from JDRF and the consensus project leadership. The initial working group comprised 61 internationally recognised physicians, nurse practitioners, clinical psychologists and DCES, with expertise in the diagnosis and care of people with early-stage type 1 diabetes. The conference was centred on monitoring of IAb⁺ people in early-stage type 1 diabetes, including discussions of current guidance on current best practice for monitoring, as applied by several prospective type 1 diabetes prevention trials (discussed in detail below).

Following a moderated discussion, expert participants were offered the opportunity to join at least one of four working groups, each focused on key aspects of monitoring. Each working group was chaired by two expert contributors, as noted below, and was tasked with self-organised review of the available evidence, participation in serial online discussions and development of core principles. The working groups were: (1) monitoring in children and adolescents (Chairs: REJB and KJG); (2) monitoring in adults (Chairs: RS-R and JMW); (3) educational needs (Chairs: KJB and BIF); and (4) psychosocial interventions (Chairs: KAD and LBS). This subsequently generated 21 separate online group discussions. Each aspect of these discussions was documented with support from JDRF team members and a medical writer. It must be noted that this document is not intended or structured as a systematic review.

On a weekly basis, from 3 May 2023 onwards, evidence-based statements and expert interpretations were drafted for review and revision. At the end of this iterative process, an agreed narrative review of the available evidence was compiled along with the expert clinical advice. Each bulleted principle was assigned a level of supporting evidence (A, B, C or E; see electronic supplementary material [ESM] Table 1) that adheres to the evidence-grading system for ‘*Standards of Care in Diabetes—2023*’, published by the ADA [75]. The process concluded with a conference to review and endorse the penultimate consensus report at the ADA’s 83rd Scientific Sessions in San Diego, CA, USA. Following this meeting, a revised draft was made available for public comment, after which the consensus document was finalised. The outcomes of this process are also summarised in an algorithm that details the decision path for monitoring of IAb⁺ people regardless of whether they were screened as part of a research protocol or in the clinical setting for any reason (Fig. 1).

1. Terminology

Precise and consistent language is important to facilitate clear communication and education. As the field has evolved, so has the language around multiple IAb⁺ status, the stages of type 1 diabetes and associated risk of progression. It was once commonplace to refer to ‘risk of’ and ‘prevention of’ type 1 diabetes in individuals with multiple IAb⁺ status. However, the staging criteria recognise seroconversion to multiple IAb⁺ status as the onset of early-stage type 1 diabetes and, thus, it is not possible to both have a condition and be ‘at risk’ for it.

Therefore, stage 1 type 1 diabetes and stage 2 type 1 diabetes (Table 1) should be referred to by their defined names or collectively referred to as ‘early-stage type 1 diabetes’. While the staging criteria are still becoming widely known, it may be appropriate to refer to these stages as ‘presymptomatic type 1 diabetes’ for some audiences, to highlight that these early stages exist prior to traditional, symptomatic (i.e. stage 3 type 1 diabetes) disease. Individuals with a genetic risk (based on genetic screening and/or family history) or with only single IAb⁺ status have pre-stage 1 type 1 diabetes and can be referred to as ‘at risk’, but individuals with multiple IAb⁺ status are confirmed as having early-stage type 1 diabetes. It must also be clear what the focus of prevention is; for example, prevention of seroconversion, progression to dysglycaemia or of stage 3 type 1 diabetes.

2. Partnership between primary-care and specialist HCPs

There is a need for primary care to take on some of the early-stage monitoring and managing IAb⁺ children and adults. However, staging criteria are relatively new and are unlikely to be widely known among primary-care HCPs. Therefore, educational steps and materials must facilitate the partnership between primary-care HCPs and secondary care. Primary-care HCPs in some regions (e.g. the USA, Europe) are involved in screening and monitoring tasks for hypercholesterolaemia and other metabolic syndromes, so the expectation is that this is possible for early-stage type 1 diabetes. A critical need is that all HCPs recognise that some IAb⁺ individuals can progress rapidly, whereas others may not develop symptoms for decades. In this context, the following expert clinical advice is suggested:

Clinical roles and responsibilities

- Primary-care HCPs should understand the stages of type 1 diabetes, as well as methods for and suggested frequency of metabolic monitoring that can be used to prevent DKA at onset of clinical type 1 diabetes. [E]
- Primary-care HCPs with a specific interest in managing people with early-stage type 1 diabetes can serve as a local referral resource for other primary-care HCPs when specialist care providers are not readily accessible. [E]
- The primary-care provider and specialty care provider, along with the at-risk single IAb⁺ individual or the multiple IAb⁺ individual with early-stage type 1 diabetes and their family, should determine which provider will have primary responsibility for metabolic monitoring and what degree of collaboration is desired. [E]
- The level of specialist engagement will need to be reassessed and may shift over time as the IAb⁺ individual progresses through the stages of type 1 diabetes, as well as when other needs and circumstances change. [E]

Communication and co-ordination of care

- Within a medical practice, HCPs should ensure that the medical record for a child, adolescent or adult, who is single or multiple IAb⁺, reflects their status and their individual plan for routine metabolic follow-up and for urgent evaluation if symptoms of hyperglycaemia develop. [E]
- If an IAb⁺ individual meets the criteria for stage 2 type 1 diabetes (Table 1), a referral should be made to a diabetologist/endocrinologist to discuss early treatment options and individualised risk of progression to clinical type 1 diabetes. [E]
- If an IAb⁺ individual develops symptomatic hyperglycaemia, an immediate consultation with, and referral to, a multidisciplinary diabetes team comprising specialists with training and expertise in diabetes is necessary. [E]

Training and skills development

- Both monitoring and education require a broader understanding of early-stage type 1 diabetes across the medical community. Inclusion of an understanding of the continuum of type 1 diabetes into all levels of medical and nursing education will require development of competencies appropriate to the role (Fig. 2). [E]

3. Monitoring in children and adolescents

The current landscape of monitoring children and adolescents in early-stage type 1 diabetes

The following section encompasses monitoring of children and adolescents aged up to 17 years. The overall algorithm is summarised in Fig. 1. For a young person who has screened positive for multiple IAb⁺ status, monitoring recommendations are also provided by the ISPAD [16] and the Fr1da study [20].

This expert clinical advice emphasises the need to benchmark the glycaemic stage of disease and to offer ongoing monitoring for disease progression, which should be appropriate to the needs of the affected person and their family. At present, standard 2 h OGTT (1.75 g of glucose per kg of body weight up to 75 g maximum) is the preferred modality, particularly for inclusion in research studies, whereas less-intensive methods are suggested for children or adolescents who decline to undertake OGTT or participate in a research protocol. Even in a clinical-study setting, adherence with OGTT monitoring can be low [76]. Given the diverse settings and resources available, amongst the monitoring tools identified (Table 5), HbA_{1c} testing is not suitable outside of the clinical setting and only random glucose assessments, routine SMBG and CGM, that do not require venipuncture, can be self-managed at home. Studies using CGM in small cohorts of children and youth with stage 1 or stage 2 type 1 diabetes have suggested that glucose levels ≥ 7.8 mmol/l (≥ 140 mg/dl) for $>10\%$ of each day is associated with an 80% risk of progression to type 1 diabetes within 12 months of the CGM assessment period [72, 77]. In this context, risk of progression to stage 3 type 1 diabetes within 2 years of baseline CGM assessment was 40% in individuals with early-stage type 1 diabetes who spent $\geq 5\%$ of each day with glucose ≥ 7.8 mmol/l (≥ 140 mg/dl) [71]. These outcomes indicate a need for more evidence to confirm the emerging value of CGM in monitoring individuals with early-stage type 1 diabetes and to understand the disease-predictive value of additional CGM metrics. This need is more pressing given that home use of CGM systems and CGM-derived glycaemic metrics is being evaluated for risk stratification for healthy relatives of people with type 1 diabetes [78, 79].

Monitoring at a 6–12 monthly cadence has been used for participants in prevention trials, depending on risk stratification. More-frequent monitoring can be indicated for children who screen positive for islet autoantibodies before 3 years of age and are at high risk of progression [24, 51], for example, at 3- to 6-monthly intervals, depending on staging [24]. It should be noted that, amongst monitoring tools, not all CGM systems are generally available in all regions, or for use in very young children. For all individuals outside of the

research setting, reducing the frequency of monitoring can be considered as part of a minimally burdensome approach and modelling studies suggest this can be achieved while meeting the goal of DKA prevention on a population level [80]. In this context, youths of Black race and/or Hispanic ethnicity are less likely to participate with monitoring [81].

Monitoring for single IAb⁺ at-risk children

Evidence from cohort studies indicates that up to 50% of children with single IAb⁺ status revert to being islet autoantibody negative (IAb⁻) [82, 83]. Children with confirmed persistent single IAb⁺ status are not at high risk for progression compared with those with multiple IAb⁺ status, with one population-based study indicating that the 10 year risk of progression to type 1 diabetes for persistent single IAb⁺ children is 14.5%, with most of that progression (10%) happening in the first 2 years after becoming IAb⁺ [51]. This analysis also showed that the progression rate is higher for young children who have single IA-2A positivity (40.5%), compared with GADA positivity (12.9%) or IAA positivity (13.1%) [51]; however, it must be noted that fewer than 10% of children with single IAb⁺ status are IA-2A⁺. Younger age (<5 years) at first single-confirmed islet autoantibody positivity is a risk factor for progression to multiple islet autoantibody positivity, particularly during the first 2 years after seroconversion [84, 85]. As children age, relative risks for progression with each antibody subtype changes [56], with an increased effect for GADA with increasing age and a reduced effect for IAA [86].

For young children, evidence indicates that metabolic and autoantibody monitoring frequency in the first 2 years after first detection of an autoantibody is key, as this is when spread from at-risk single islet autoantibody positivity to early-stage type 1 diabetes with multiple islet autoantibody positivity is most likely. Following confirmed single IAb⁺ status, the IAb⁺ evolution after 2 years predicts development of clinical type 1 diabetes [87]. Progression to multiple IAb⁺ status or reversion is also highest in the first 2 years in single IAb⁺ pre-school children, with a hazard rate of 0.3 in the first 2 years vs 0.05 for children who have been single IAb⁺ for >2 years [84]. Among children with increased genetic risk, those who remain single IAb⁺ have a risk for type 1 diabetes of 1.8 per 100 person-years, children who revert to negative status have a risk of 0.14 per 100 person-years and children who have never been IAb⁺ have a risk of 0.06 per 100 person-years [83]. The rate of progression to multiple IAb⁺ status also declines with age [88].

Expert clinical advice for monitoring of single IAb⁺ (at-risk) children

- Confirm persistent single IAb⁺ status after first detection in a second sample, preferably in a laboratory that meets IASP standards, using two independent methods [89], and confirm negative status for other islet autoantibodies. **[B]**
- Islet autoantibody status and metabolic monitoring during the first 2 years after seroconversion is most critical [51, 84, 85, 90]. Ongoing metabolic monitoring is not essential beyond this 2 year period. **[B]**
- Children who develop type 1 diabetes at a very young age have more-rapidly progressing and aggressive disease. For children aged <3 years who are single IAb⁺, monitor their IAb⁺ status every 6 months for 3 years, then annually thereafter for 3 more years. Metabolic monitoring in children aged <3 years should include random venous or capillary blood glucose and HbA_{1c} values at the same frequency [51, 84, 85, 87, 90]. If no progression, stop autoantibody and metabolic monitoring, and counsel for risk of clinical disease. **[B]**
- For children aged ≥3 years at first positive test, monitor IAb⁺ status annually for 3 years. Metabolic monitoring should include annual random venous or capillary blood glucose and HbA_{1c} testing for 3 years [51, 84, 85, 90]. If no progression after 3 years, stop autoantibody and metabolic monitoring, and counsel for risk of clinical disease [51, 84, 85, 90]. **[C]**
- For children with single islet autoantibody positivity who revert to seronegative during autoantibody monitoring, or do not progress (see above), education should be provided to their families emphasising potential symptoms and awareness of DKA [33, 91, 92]. **[C]**

Limitation Many data on single IAb⁺ children are derived from groups with extended prospective follow-up and known genetic risk profiles or first-degree relatives with type 1 diabetes with limited racial/ethnic diversity. Data on individuals in the general population are more limited, particularly in those with a single screening event.

Monitoring for multiple autoantibody-positive children (early-stage type 1 diabetes)

Children with confirmed multiple IAb⁺ status are at very high risk for progression to stage 3 type 1 diabetes within 15 years. Combined data from five prospective studies indicate that the 15 year risk for stage 3 type 1 diabetes is 85% for children with two islet autoantibodies and 92% for those with three islet autoantibodies, and that there is a >99% lifetime risk [87]. In children with multiple islet autoantibody positivity, younger age at first islet autoantibody detection predicts more-rapid progression to stage 3 type 1 diabetes [51, 93]. Although data on children with multiple islet autoantibody positivity identified from general population screening are derived from shorter follow-up durations, progression rates appear to be similar to those observed in relatives of individuals with type 1 diabetes enrolled in longitudinal research cohort studies [24, 94].

The detection of multiple autoantibodies should be confirmed in a venous sample, within 3 months [49]. However, this should not be a rate-limiting step in the monitoring or treatment process, as progression can happen rapidly in young children. Confirmation is critical, since without it there is a risk of delivering a false diagnosis of multiple IAb⁺ status, with consequent anxiety and distress for the individual. Conversely, although loss of confirmed multiple IAb⁺ status is rare and may be associated with reduced risk of progression to type 1 diabetes [95], monitoring should not be discontinued in this group.

Expert clinical advice for monitoring of multiple IAb⁺ children (early-stage type 1 diabetes)

Monitoring of glucose metabolism among children with multiple IAb⁺ status is necessary to predict time to stage 3 diagnosis, identify those who may be eligible for intervention and prevent DKA. Options for metabolic assessments include home SMBG monitoring, periodic CGM assessment, and laboratory testing for HbA_{1c}, random venous or capillary blood glucose and OGTT (with stimulated C-peptide assessments). It is acknowledged that there is variable access to high-quality laboratory-testing facilities outside of the research setting. Where possible, the opportunity to undertake monitoring at home or in the primary-care setting should be considered (Table 5).

Expert clinical advice for monitoring of multiple IAb⁺ children (early-stage type 1 diabetes)

- Education must be provided to reinforce the need for and value of longitudinal monitoring to prevent DKA [33, 91, 92]. Written instructions with relevant emergency contact details should be provided in case of type 1 diabetes symptoms and/or hyperglycaemia. **[E]**
- Confirm persistent multiple IAb⁺ status after first detection in a second sample, preferably in a laboratory that meets IASP standards, following the 'rule of twos' [52], preferably using two independent methods [89]. **[B]** Where a two-test confirmation is not possible, a single blood test positive for multiple islet autoantibody status identifies a person with sufficient risk for metabolic monitoring. **[E]**
- In infrequent cases, for a child with previously confirmed multiple IAb⁺ status and who has reverted to single IAb⁺ or IAb⁻ status [95], monitoring should also follow the advice below. **[E]**
- Metabolic monitoring should be conducted based on the staging criteria and modalities described in Table 1 and Table 5. This should be undertaken when the child is healthy and not experiencing intercurrent illness. **[E]**
- SMBG meters and strips can be provided to all children with multiple islet autoantibody positivity or their parents. **[E]**
- During intercurrent illness, SMBG can be used to detect hyperglycaemia. **[A]**
- For children with recent confirmation of multiple IAb⁺ status, an SMBG test can be performed on two different days over a 2 week period (on each day, test either fasting or postprandial), and again, thereafter, once every 1–3 months. See also advice below. **[E]**
- In children with stage 1 type 1 diabetes, HbA_{1c} should be measured once every 3 months for children <3 years old, at least every 6 months for children 3–9 years old and at least every 12 months for children >9 years old [93]. **[E]** Increase in longitudinal HbA_{1c} of ≥10%, even in the normal range (e.g. from 31 mmol/mol [5.0%] to 37 mmol/mol [5.5%]) indicates increased risk of disease progression to stage 3 type 1 diabetes within a median of 1 year [66, 67]. **[B]**
- In children with stage 2 type 1 diabetes, measures of glucose regulation should be monitored every 3 months, as above. **[E]**
- Longitudinal change in HbA_{1c} of ≥10% from date of confirmed islet autoantibodies may indicate dysglycaemia and disease progression [66, 67], and requires the performance of an OGTT to assess type 1 diabetes stage (Table 1) in order to determine eligibility for therapy. **[E]**
- Random venous or capillary blood glucose should be measured at the same time as HbA_{1c}. Rise in venous glucose in children with multiple IAb⁺ status predicts time to stage 3 type 1 diabetes (see Table 1) [96]. **[E]**
- OGTT is the established gold standard to classify stage 1, stage 2 or stage 3 type 1 diabetes **[A]**, but if performing OGTT is not possible, obtain a 2 h postprandial capillary blood glucose after a carbohydrate-rich meal to assess for dysglycaemia [86]. **[E]**
- Monitor objective weight trends in a growing child using a growth chart, **[C]** which may be below the normal range during progression of type 1 diabetes. Ensure that a healthy meal plan has been maintained to preclude disordered eating behaviours as a cause of weight change. **[E]**
- 10–14 day CGM can be used periodically to monitor glucose metabolism at a similar frequency as HbA_{1c} measurement. **[E]** CGM should ideally be blinded to the individual wearing it and must be interpreted by trained HCPs, with education for the user and their family. **[E]** Criteria for CGM metrics to diagnose stage 2 or stage 3 type 1 diabetes are proposed (Table 1) and require further research.
- Stage 2 type 1 diabetes warrants referral to specialists in type 1 diabetes progression for discussion of risk and options for monitoring, wherever feasible. **[E]**
- In countries with approved therapeutic options for early-stage type 1 diabetes or locations with access to intervention studies [3, 97], referral to a clinical centre with expertise in the specific treatment should be done when stage 2 type 1 diabetes is suspected or diagnosed. **[E]**

4. Monitoring in adults

The current landscape of monitoring adults who are at risk of or have early-stage type 1 diabetes

The following guidance encompasses monitoring of adults aged 18 years and over, although the advice is based on outcome data that typically reflect adults younger than 45 years of age. Data specific to adults older than this are an important unmet need. Epidemiological data show that, overall, type 1 diabetes is diagnosed more frequently in adulthood than in childhood [98–101], at a median of more than 35 years of age [102, 103]. Despite this, misdiagnoses of type 1 diabetes in adults remain common and are increasingly likely with age [60], setting the scene for development of DKA. In common with childhood-onset type 1 diabetes, adult-onset type 1 diabetes is associated with the presence of islet-specific autoantibodies [104–107]. Although TrialNet cohort data indicate that the rate of progression to type 1 diabetes in IAb⁺ adults is slower than in children, many adults with multiple IAb⁺ status and early-stage type 1 diabetes still develop stage 3 disease [59]. While it has been suggested that progression in some adults may not occur and that some of those who do progress have only single islet autoantibody positivity, further long-term follow-up data are needed to better characterise the long-term implications of persistent autoimmunity in adults [108]. For example, recent data highlight the frequent presence of islet autoimmunity in cohorts presenting with phenotypic type 2 diabetes [109].

Guidance to inform clinical monitoring practices in adults represents a considerable unmet need. There are many evidence-base gaps, including a lack of information about risk of disease progression in IAb⁺ adults without a family history of type 1 diabetes, particularly in individuals with non-European ancestry. Data on suggested monitoring protocols, including effectiveness in preventing DKA and adherence with monitoring, are substantially based on children and adolescents. The frequency of DKA among adults at diagnosis with type 1 diabetes is unknown but believed to be lower than for children, given that adults may recognise and respond to symptoms of hyperglycaemia, and often have higher C-peptide levels at clinical diagnosis and a slower decline in beta cell function over time [110]. Yet, incorrect assumptions leading to underdiagnosis of type 1 diabetes in adults mean many develop DKA before starting insulin therapy.

DKA incidence at clinical diagnosis can be reduced by participation in active monitoring [24, 26, 27]. Regarding frequency of monitoring, modelling based on TrialNet data suggest that conducting approximately half the number of visits involved in a research setting (typically once every

12 months rather than every 6 months), is likely to be effective in substantially reducing the incidence of DKA to the levels seen in research studies both for children and adults [80]. However, data from the TrialNet study indicate that adults 18 years and older are less likely than paediatric participants to engage with recommended monitoring using 6–12 monthly OGTT in the early phases after screening positive for autoantibodies [81]. As with youths, adults of Black race and/or Hispanic ethnicity are less likely to participate with monitoring in this context [81].

Most endocrinologists and primary-care HCPs will not be trained in monitoring adults with single IAb⁺ status or early-stage type 1 diabetes. Thus, the educational need will be significant. As with children and adolescents, monitoring in IAb⁺ adults must be realistic and actionable across diverse regions, with different resources. HCPs are significantly burdened such that additional tasks for monitoring in pre-stage 3 type 1 diabetes must be clinically useful.

Expert clinical advice for monitoring single IAb⁺ (at-risk) adults

- Confirm persistent single IAb⁺ status after first detection in a second sample, preferably in a laboratory that meets IASP standards, using two independent methods [89], and confirm negative status for other islet autoantibodies. **[B]**
- Annual metabolic monitoring should be considered for single IAb⁺ adults if there are additional risk factors, including one or more of the following: first-degree relative with type 1 diabetes; elevated genetic risk for type 1 diabetes if tested; dysglycaemia (e.g. impaired fasting glucose or impaired glucose tolerance); or history of stress hyperglycaemia [111, 112]. **[E]**
- Although single IAb⁺ adults are at lower risk of progression to type 1 diabetes compared with children [59], and this risk continues to fall with increasing age, there remains a residual risk for progression. The approach to metabolic monitoring for single IAb⁺ adults can be informed by that applied for screening for type 2 diabetes, which is advised every 3 years for normoglycaemic adults aged >35 years, or who are overweight/obese with one or more additional risk factors [18]. A similar 3 year frequency is proposed for single IAb⁺ adults, to monitor for risk of progression, which may be increased to annual monitoring with the additional risk factors identified for type 2 diabetes. **[E]**
- No type 1 diabetes monitoring is indicated in individuals with transient single islet autoantibody positivity who then revert to being seronegative. Screening for diabetes in this group of adults should, thereafter, follow standard of care guidelines for type 2 diabetes [18]. **[C]**

Monitoring for single autoantibody-positive at-risk adults

Frequency of monitoring can be based on the stage at which an individual with islet autoantibody positivity is diagnosed. Single IAb⁺ adults with dysglycaemia should be monitored more frequently than those with normoglycaemia. Additional risk stratification may also be possible based on other characteristics, such as age, or modifiable factors, such as abdominal obesity.

Monitoring for multiple autoantibody-positive adults (early-stage type 1 diabetes)

As with monitoring in single IAb⁺ adults, more-frequent monitoring is proposed for individuals with multiple IAb⁺ status if they are diagnosed with stage 2 type 1 diabetes compared with stage 1 type 1 diabetes. Risk stratification based on age, abdominal obesity and other modifiable factors also applies.

Expert clinical advice for monitoring multiple IAb⁺ (early-stage type 1 diabetes) adults

- Education must be provided to reinforce the need for and value of longitudinal monitoring to prevent DKA [33, 91, 92]. Written instructions with relevant emergency contact details should be provided in case of type 1 diabetes symptoms and/or hyperglycaemia. **[E]**
- Confirm persistent multiple IAb⁺ status after first detection in a second sample, following the ‘rule of twos’ [52] and preferably using two independent methods [89]. **[B]** Where a two-test confirmation is not possible, a single blood test positive for multiple islet autoantibody status identifies a person with sufficient risk for metabolic monitoring. **[B]**
- In infrequent cases, for adults with previously confirmed multiple IAb⁺ status who have reverted to single IAb⁺ or IAb⁻ status [95], monitoring should also follow the advice below. **[E]**
- All multiple IAb⁺ adults can be provided with SMBG meters and strips to be used during illness or when symptoms may be present. **[E]**
- In adults with stage 1 type 1 diabetes and normoglycaemia (Table 1), glycaemic status should be monitored using HbA_{1c} every 12 months, as part of routine primary-care visits. Modify frequency of monitoring based on individual risk assessment, based on age, number and type of islet autoantibodies, and glycaemic metrics [4]. **[E]**
- If duration of normoglycaemia extends to 5 years, metabolic monitoring every 2 years may be sufficient. **[E]**
- In adults with confirmed stage 2 type 1 diabetes (Table 1), metabolic status should be monitored using HbA_{1c} every 6 months, in conjunction with one other of the following monitoring modalities: blinded CGM (applied and interpreted by trained HCP); higher frequency of SMBG; or 2 h plasma glucose following 75 g OGTT. **[E]**
- Longitudinal change in HbA_{1c} of ≥10% from date of confirmed islet autoantibodies may indicate dysglycaemia and disease progression [66, 67], and requires the performance of an OGTT to assess type 1 diabetes stage (Table 1) in order to determine eligibility for therapy. **[E]**
- When dysglycaemia or hyperglycaemia occurs, C-peptide monitoring should be considered where the diagnosis of type 1 diabetes vs type 2 diabetes is unclear. Meta-analysis indicates that a C-peptide level of ≤0.20 nmol/l with IAb⁺ status can be associated with a diagnosis of type 1 diabetes rather than type 2 diabetes [113]; however, many adults presenting with type 1 diabetes will have C-peptide above this level [110]. **[B]** Note that C-peptide levels can be falsely low in hypoglycaemia (<3.9 mmol/l [<70 mg/dl]), after fasting or in severe hyperglycaemia/DKA, so concomitant plasma glucose concentration should be checked and interpreted in combination with the clinical state.
- In countries with approved therapeutic options for early-stage type 1 diabetes or locations with access to intervention studies [3, 96], referral to a clinical centre with expertise in the specific treatment should be done when stage 2 type 1 diabetes is suspected or diagnosed. **[E]**

Monitoring during pregnancy for IAb⁺ women

Evidence on the progression of type 1 diabetes in IAb⁺ pregnant women is limited and research data on this aspect of managing risk in early-stage type 1 diabetes is a significant unmet need (Table 6). With that said, a high risk for postpartum type 1 diabetes has been indicated [114], and the guidance below is primarily based on expert opinion. Pregnancy demands increased pancreatic beta cell function and may result in diabetes, as it does in gestational diabetes mellitus (GDM) [115]. Given that 60% of babies born to women with diagnosed type 1 diabetes are large for gestational age (LGA), which is associated with increased rates of obstetric and neonatal complications [116, 117], it is important to avoid a missed early diagnosis and promote normal fetal development.

Expert clinical advice for monitoring in pregnancy for IAb⁺ women

- Women with confirmed islet autoantibody positivity who become pregnant should have an OGTT, HbA_{1c} test or application of CGM soon after pregnancy is confirmed (by 8 weeks if possible) [18, 118]. [C]
- Women with confirmed islet autoantibody positivity who are not already diagnosed with type 1 diabetes should receive OGTT tests at 24–28 weeks of pregnancy, as standard for all pregnancies [18]. [A]
- Glucose monitoring for women with confirmed IAb⁺ status who are diagnosed with type 1 diabetes: once postpartum, women should be assessed prior to discharge from hospital, in consultation with a specialist endocrinologist, to determine continued need for insulin [114]. [C]
- Women with confirmed IAb⁺ status should be monitored for 6–12 months postpartum to assess any changes in insulin requirement. [E] Where available, follow-up both with the gestational care provider and an insulin-initiation specialist should be provided. [E]

5. When to start insulin

At some point, monitoring will reveal a person with persistent and/or recurrent hyperglycaemia prompting a decision on whether to start insulin, along with associated education and support for affected individuals and their families. As screening programmes identify more people with early-stage type 1 diabetes, more people are being assessed as meeting classic diagnostic criteria for stage 3 type 1 diabetes (Table 1), but who might not yet require insulin therapy. Decisions about how and when to initiate insulin will be based on a range of factors, many of which

do not have a body of evidence. Therefore, consideration of starting insulin should trigger a referral to a specialist centre with expertise in initiating and managing people with type 1 diabetes on insulin.

6. Education

The primary goals of education for the care of IAb⁺ individuals and their families are outlined in Table 7. Given the paucity of evidence on education for people with early-stage type 1 diabetes, extensive experience in education for stage 3 type 1 diabetes can be extrapolated to this population. National standards for diabetes self-management education and support (DSMES) have been published by the ADA and the Association of Diabetes Care & Education Specialists (ADCES) and are broadly applicable in this context [119]. When appropriate, evidence from studies in stage 3 type 1 diabetes are used to support grading of evidence.

Experience in clinical studies can also inform education for people with early-stage type 1 diabetes and their families/caregivers. The Environmental Determinants of Diabetes in the Young (TEDDY) prospective study protocol emphasises parental education regarding symptoms and signs of diabetes. For families new to type 1 diabetes, this education provides foundational skills for diabetes management that are a component of reduced parenting stress at the time of stage 3 diagnosis compared with individuals who were members of the community control group and did not receive education [120]. Similarly, families of children with early-stage type 1 diabetes in the Fr1da study are invited to participate in an educational programme of blood glucose monitoring and symptoms of hyperglycaemia/DKA. They are also provided with a guidebook specifically designed for children with early-stage type 1 diabetes, and assigned a contact person to answer questions at any time. Children who take part in this programme alongside metabolic monitoring have a lower rate of DKA and reduced HbA_{1c} at stage 3 type 1 diabetes presentation compared with children who declined education and follow-up [33]. Over 50% had no symptoms at the clinical presentation of stage 3 type 1 diabetes, 93.5% had no weight loss and length of stay in hospital was shorter [91, 92].

Basic community awareness campaigns not associated with monitoring and centred on the early symptoms of type 1 diabetes, that target teachers, paediatricians and parents, have been effective in reducing DKA rates in children in regional settings (Parma in Italy [121] and Newcastle in NSW, Australia [122]). However, national campaigns in Italy and Austria, with the same objectives, have not seen the same impact [123, 124]. The content and delivery of these campaigns were not similar, so it is hard to draw conclusions about the effectiveness of this education.

Table 6 Selected unmet needs for further research and clinical development

Unmet research needs

- Long-term rates of progression to stage 3 diabetes in IAb⁺ individuals without a family history of T1D, and progression rates in adults and people of non-European ancestry.
- The impact of pregnancy in women who are IAb⁺ and the glycaemic changes that may be evident during pregnancy and in the postpartum period, along with risks for progression to stage 3 T1D during and after pregnancy.
- Neonatal outcomes for infants of women who are IAb⁺ and the association with glycaemic changes during pregnancy.
- Cost-effectiveness of monitoring strategies for individuals with early-stage T1D.
- Timing of insulin initiation in people with presymptomatic T1D, including short- and long-term metabolic and mental-health outcomes of different strategies.
- Impact of education alone, independent of other monitoring activities, on frequency of DKA at diagnosis and presentation of T1D.
- Methods of identifying and monitoring behavioural health needs in early-stage T1D.

Unmet clinical needs

- Comprehensive and consistent educational materials that use consistent language and vocabulary when referring to diabetes stages and risk, including translation into region-specific languages. This applies to all impacted people, from affected individuals to expert providers.
- Validated tools to measure the anxiety, depression and other mental-health behaviours that are specific to early-stage T1D.
- Sufficient availability of mental-health professionals with expertise in T1D, including early-stage T1D in youth and adults.
- Knowledge and coverage of appropriate monitoring by stakeholders (insurers, clinicians, etc.)
- Timely access to expert HCPs and centres of expertise for intervention(s) to delay onset of stage 3 T1D.

The key principles presented in this table and in this consensus document will be subject to updating once additional evidence becomes available T1D, type 1 diabetes

Education topics and intensity for people with early-stage type 1 diabetes and their families should be based on type 1 diabetes stage, age, rate of progression, etc. First-degree relatives may have different needs for support and guidance from the general population, as they have an established awareness of the implications and impact of IAb⁺ status. Education topics should be linked to specifically timed action plans and include the topics detailed below. Education can be tailored so it is uniquely appropriate for both stage 1 and stage 2 type 1 diabetes (Table 1). Clinical practitioners with experience in early-stage type 1 diabetes should be involved in the later steps of education.

When should education be provided?

The needs for education are centred on the key moments in the life of the person with early-stage type 1 diabetes [119].

These are at the point of a positive autoantibody screen, at diagnosis of each stage, when monitoring tasks are performed, and annually for review and maintenance. Education is also critical during life transitions and milestones, and when care needs change.

Key education topics

Education and self-care behaviours for individuals at risk for or with early-stage diabetes (Table 1) can be derived from the overall framework of self-management skills for diabetes and related conditions. These are described in the ADCES7 self-care behaviours [125]. Those relevant for at-risk individuals or those with early-stage type 1 diabetes focus on understanding the implications of their single (at-risk) or multiple (early-stage type 1 diabetes) IAb⁺ status, and the benefits of regular monitoring. Symptom awareness and

Table 7 The primary goals of education for care of IAb⁺ individuals and their families

1. To prevent DKA and promote safe monitoring practices and reduce the occurrence of symptoms of diabetes
2. To minimise the requirement for emergency care, hospital admission and need for intensive care at diagnosis of T1D
3. To improve appropriate risk perception at each monitoring milestone
4. To understand specific outcomes, e.g. prevention of DKA, initiation of insulin therapy
5. To understand available interventions
6. To explore and understand the benefits of individual participation in research studies
7. To provide education that supports psychosocial interventions to optimise general health and mental health for affected individuals and their families

T1D, type 1 diabetes

metabolic monitoring are important to reduce the risks of hospitalisation for DKA. If other family members have type 1 diabetes, HCPs should not assume pre-existing awareness and knowledge. The most-current education should always be mandated.

Educational topics of highest value for IAb⁺ individuals and family members

For an individual who has tested positive for one or more autoantibodies, a person-centred plan should be developed that is best suited to the IAb⁺ person and their individual situation. Their family members should be included as part of the programme of education. The topics that may have high value are likely to include the following: (1) understanding autoimmunity and the confirmation of single (at-risk) or multiple (early-stage type 1 diabetes) IAb⁺ status; (2) definition of at-risk or early-stage type 1 diabetes; (3) risk perception (accurate risk perception is linked with staying engaged in monitoring and with DKA prevention [126]); (4) risks and benefits of individual participation in research studies; (5) awareness of hyperglycaemia episodes for introducing insulin at the right time; (6) strategies for healthy coping; (7) symptom awareness and prevention of DKA; (8) glucose monitoring (SMBG, CGM), if clinically recommended; (9) healthy behaviours, including meal planning and physical activity; (10) risks and benefits of intervention therapies; (11) monitoring planning, with descriptions of laboratory tests and devices that may be used (Table 5); and (12) treatment options and introduction to insulin therapy.

Where should education be provided?

Education should be widely accessible via a variety of modalities, across multiple media platforms and settings, and should be crafted with the specific audience's learning needs in mind. For education aimed at HCPs, a key requirement is for professional associations in all regions to be aligned with the educational programme and curriculum, preferably compatible with their educational platforms and with accreditation. For education aimed at people with pre-stage 3 type 1 diabetes, in-person options associated with clinical appointments or in-group sessions are important, and strong evidence supports DSMES delivery through virtual, telehealth, telephone, text messaging and web-based/mobile phone applications (apps) [127–131].

Who should provide education?

The competencies that must be addressed in education are outlined in Fig. 2. There is a need for diabetes professional associations to endorse the educational goals, educational

tools and educational content, as described. Different groups of individuals, including HCPs, community members, and individuals in need of monitoring and their families (indicated in the pyramid sections in Fig. 2), should have the competencies described and participate as appropriate.

Expert clinical advice for education of single IAb⁺ (at-risk) and multiple IAb⁺ (early-stage type 1 diabetes) individuals

- Education is the responsibility of all health professionals involved in the monitoring and care of individuals with type 1 diabetes. [E]
- People who are at risk or with early-stage type 1 diabetes may participate in monitoring education programmes to reduce the rate of DKA at diagnosis [33, 90, 91]. [B]
- Education should be provided: (1) at the point of a positive autoantibody screening; (2) at diagnosis of each stage; (3) annually for review and maintenance; and (4) during life transitions. [E]
- Education should accompany the implementation of all monitoring plans; this includes home glucose testing and any monitoring devices. [E]
- Education should be culturally, linguistically and socioeconomically congruent. [E]
- Education topics and intensity should be based on type 1 diabetes stage and risk of progression, and include the risks and benefits of intervention therapies, when appropriate. [E]
- Diabetes education should be accessible, engaging and person-centred. This includes consideration of the developmental, social, emotional, cultural and linguistic needs of the individual and/or their family. [E]

7. Psychosocial support

What is the current landscape regarding psychosocial support for people with type 1 diabetes-related autoantibodies?

People who learn that they or a loved one have type 1 diabetes-related autoantibodies often experience significant stress [132]. This is in part because events that are unpredictable, uncontrollable and threatening may be highly stressful. People who have islet autoantibody positivity, particularly those who have multiple islet autoantibody positivity, will very likely develop type 1 diabetes in the future. However, disease progression is impossible to predict precisely and having IAb⁺ status does not mean imminent type 1 diabetes onset [133, 134]; stage 3 type 1 diabetes, with associated insulin

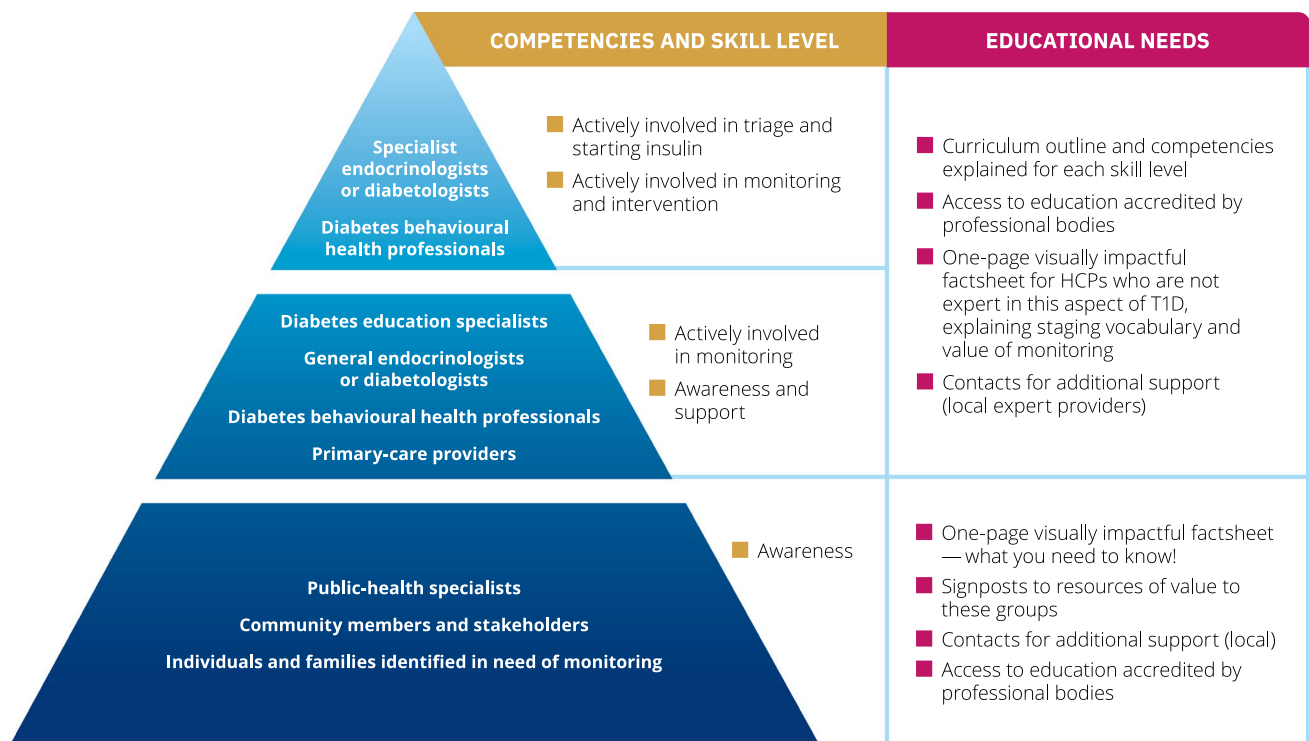


Fig. 2 The continuum of educational needs and competencies: what does one need to know? The image represents the anticipated skills that must be developed within the continuum of stakeholders in monitoring presymptomatic type 1 diabetes. The groups indicated within

the pyramid sections should have the competencies described and participate as appropriate. The need is for unified, consistent, globally applicable language at all levels. T1D, type 1 diabetes. This figure is available as part of a [downloadable slideset](#)

administration and glycaemic monitoring, could be months or even years away [17].

When learning they have type 1 diabetes-related autoantibodies, individuals of all ages and their family members can experience a range of emotional and behavioural reactions [135, 136], including shock, grief, guilt, anger, depression and anxiety. If time passes with no diagnosis of stage 3 type 1 diabetes, cognitions about type 1 diabetes may change and individuals may become convinced they will never get the disease or have reduced risk, despite evidence to the contrary [137]. Parents often engage in behaviours in attempts to prevent type 1 diabetes when faced with the news that their child is at increased risk, even when not provided with recommendations to do so, though more-recent data have shown that lower physical activity and meal plans with a higher glycaemic index are associated with faster progression to type 1 diabetes [138–140]. Meal-planning changes are most commonly reported, with extra monitoring at home (including blood glucose checking) being particularly common in families with someone who already has type 1 diabetes [141, 142].

Research has documented the psychosocial impact of newborn screening [143], as well as genetic and islet autoantibody screening for type 1 diabetes [132, 136]. Failure to

understand the screening and risk information presented is common. For example, more than a third of participating mothers and over half of participating fathers in the TEDDY study stated that their child was not at increased risk for type 1 diabetes, despite being clearly informed of their child's increased genetic risk [137]. To date, no data are available on how children screened positive for islet autoantibodies perceive or react to their risk.

Emotional distress in response to a positive islet autoantibody screen is also common. Many parents of children in the TEDDY study experienced anxiety after learning that their child was at increased risk for developing type 1 diabetes, with mothers reporting higher anxiety than fathers [132]. Although anxiety decreased across time for parents of IAb⁺ children who never developed additional autoantibodies, anxiety remained elevated in many parents of children with multiple autoantibodies for years after the child's first IAb⁺ test result. Mothers who experienced negative interpersonal life events and postpartum depression, but who were accurate about their child's type 1 diabetes risk, were particularly vulnerable to heightened anxiety [144]. In the Autoimmunity Screening for Kids (ASK) study, which conducted islet autoantibody screening in the general population, 74.4% of parents reported significant levels of anxiety about their

child's type 1 diabetes risk at the first follow-up visit; parents with lower educational attainment were more likely to exhibit higher levels of anxiety [145].

Around 40% of mothers and 20% of fathers in the Fr1da study reported clinically elevated symptoms of depression after learning that their child was at increased risk for type 1 diabetes as compared with around 18% of mothers and fathers of children who were IAb⁻ [24]. Depressive symptoms declined across 1 year, with scores in mothers of IAb⁺ children remaining slightly elevated as compared with mothers of IAb⁻ children; scores in fathers did not remain elevated.

Although both the ADA and ISPAD have published recommendations about the psychosocial care of individuals with stage 3 type 1 diabetes [146–148], these are limited to general principles for care of those with early-stage type 1 diabetes [149]. Thus, there is an urgent need to provide guidance on psychosocial support for individuals with type 1 diabetes-related autoantibodies and their families.

We recognise regional differences in healthcare resources may limit mental-health resources for care of people with diabetes. In most areas, there are insufficient mental and behavioural health professionals with expertise in the psychosocial aspects of type 1 diabetes who can provide the care recommended by the ADA and ISPAD [146–148].

What is the purpose of psychosocial support?

The overall goal of providing psychosocial support for individuals identified as having early-stage type 1 diabetes and their families is to assist them in successfully managing the psychosocial impacts associated with this life-changing news. To accomplish this goal, emotional, cognitive and behavioural functioning need to be assessed and addressed, not only in individuals with type 1 diabetes-related autoantibodies but in their family members as well, when appropriate.

What type of support should be provided?

The essential first step is to ask the individual who is at risk for type 1 diabetes and/or their caregivers and family members about their reactions upon receiving the news that they have type 1 diabetes-related autoantibodies. However, asking once is not enough as adjustment to autoantibody status may change over time [132]. Enquiring about how individuals are coping with the news and their current needs should be conducted at every monitoring visit. Examples of questions to include in the conversation include:

1. How do you feel about this news?
2. Others have said this news brings feelings of sadness or worry, what are your feelings?
3. What is your understanding about having multiple autoantibodies?
4. What type of things are you doing to try to prevent type 1 diabetes?
5. What are your thoughts about talking with a counsellor about your feelings from this news?

Providers can also assess global symptoms of anxiety and depression using age-appropriate standardised and validated questionnaires, such as the Patient Health Questionnaire-9 (PHQ-9) for depression [150] or the Hamilton Anxiety Scale [151]. However, global measures of anxiety and depression may not be sensitive to the emotional impact specifically associated with learning that one—or a loved one—has type 1 diabetes autoantibodies. In such cases, measures that assess emotional reactions to the IAb⁺ status, such as the 'State' component of the State-Trait Anxiety Inventory [152] may be more appropriate. At a minimum, HCPs should have conversations with individuals about their reactions to IAb⁺ results rather than relying solely on global measures of psychosocial functioning. Assessments should occur at regular intervals, since reactions are likely to change over time. Additional measures for both depression and anxiety in diabetes are provided in the ADA Psychosocial Care for People With Diabetes Position Statement [147], along with a directory of mental-health providers (<https://my.diabetes.org/health-directory>) [153]).

It is also important to consider developmental and family-specific factors when assessing psychosocial needs. For example, children and adolescents with type 1 diabetes autoantibodies may experience varied emotional, cognitive and behavioural impacts as they develop. This further emphasises the need for ongoing, regular assessment of psychosocial needs. Additionally, individuals with a family history of type 1 diabetes may react differently to learning about type 1 diabetes-related autoantibodies [141] compared with those who are unfamiliar with the disease; family context and prior experience with type 1 diabetes are important considerations when assessing psychosocial impact and the need for additional support.

Although increased anxiety and depression can occur in individuals with type 1 diabetes-related autoantibodies and their family members, this can be reduced by monitoring for the potential development of type 1 diabetes [120]. Providing individuals with regular monitoring for type 1 diabetes, depending on stage, as outlined in earlier sections of this statement, can help individuals manage some of the unpredictability of type 1 diabetes development [120, 132].

Based on the extant literature, diabetes-focused organisations, such as the ADA and ISPAD, have provided recommendations on the importance of individuals with diagnosed type 1 diabetes receiving psychosocial care [146–148] that is preferably integrated into routine diabetes visits and delivered by providers with diabetes-specific training [154]. While the same level of evidence does not yet exist in those individuals with type 1 diabetes-related autoantibodies, the well-documented emotional, cognitive and behavioural impacts of autoantibody status certainly suggest that a similar standard for psychosocial care should be available for all individuals who are at risk for developing type 1 diabetes and their families. For individuals with early-stage type 1 diabetes and their family members, there are well-developed models of managing psychosocial reactions to risk status, including age-specific education and assigned contact people to answer questions, who can also serve as role models [9, 145].

Ideally, psychosocial care should be integrated with routine monitoring visits and delivered by HCPs using a collaborative, person-centred, culturally informed approach. When available, refer to mental and behavioural health professionals with expertise in type 1 diabetes for additional assessment and treatment. For individuals residing in the USA, the ADA publishes the Mental Health Provider Directory, which lists providers with expertise in diabetes [153].

Expert clinical advice for psychosocial support for single IAb⁺ (at-risk) and multiple IAb⁺ (early-stage type 1 diabetes) individuals

- Emotional, cognitive and behavioural functioning should be assessed in people at risk or with early-stage type 1 diabetes and their family members, when appropriate. Anxiety, risk perception and behaviour changes should specifically be assessed. [E]
- As an essential first step to providing psychosocial support, HCPs should ask the individual at risk or with early-stage type 1 diabetes and/or their caregivers and family members about their reactions upon receiving the news that they have type 1 diabetes-related autoantibodies. This can be accomplished using guiding questions and standardised and validated questionnaires. [E]
- At each monitoring visit, there should be enquiries into current needs, particularly coping. [E]
- Psychological care should be integrated into routine medical visits and, whenever possible, delivered by providers with diabetes-specific training. [E]

8. Unmet needs for further research

This consensus document for monitoring individuals with single (at-risk) and multiple (early-stage type 1 diabetes) islet autoantibody positivity covers key principles based on existing evidence and agreed expert opinion. It also highlights the significant unmet need for further research on early-stage type 1 diabetes to further increase the rigour for future guidance and recommendations, and drive the evolution of clinical care for people who have tested positive for islet autoantibodies. The key principles in this consensus document will be subject to updating once additional evidence becomes available, as indicated in Table 6.

Supplementary Information The online version contains peer-reviewed but unedited supplementary material including a slideset of the figures for download, available at <https://doi.org/10.1007/s00125-024-06205-5>.

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