









DIABETES

A guide for general practice clinical management in type 2 diabetes

- Taranaki -

3rd Edition July 2024

Our aim is that a person with type 2 diabetes can go to any general practice in Taranaki and receive excellent care for their diabetes.

This folder is to give all general practice teams a resource for type 2 diabetes. It is filled with practical information, hints and tips that are intended to support people who are beginning to work in diabetes through to proficient nurses developing knowledge and clinical reasoning in diabetes care.

We have focused on clinical management; however we acknowledge there is much more to working with people who live with any long-term condition than simply clinical knowledge.

This is not an original work, but a compilation from many different sources and experts from our local area. The original manual was created in 2018, and this is the second update since then.

We hope you find it useful, and we always welcome feedback.

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The clinical diabetes specialist for the Taranaki region is Lauren Southgate. If your practice requires any support, call Lauren who will come and work alongside your team to upskill them to gain confidence in diabetes management.

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SCREENING FOR DIABETES

Adults

New Zealand Society for the Study of Diabetes (NZSSD) has recommended that HbA1c should be the first line check for screening and diagnosis of type 2 diabetes in adults.

Screening for Diabetes should be offered:

- As part of cardiovascular risk assessment (CVA)
 - In Māori, Pasifika or Indo-Asian screen men from 30 years and women from 40 years of age
 - For Europeans, screen men from 45 years and women from 55 years of age
- Those who are at high risk of developing diabetes.
- Whanau of current patients with diabetes use the PMS to locate high risk whanau.

<u>Identified risk factors for diabetes:</u>

- Non-European ethnicity
- Family history of Diabetes, including first degree relative <40 years of age
- Pre-Diabetes (HbA1c 41-49mmol/L)
- History of Gestational Diabetes or large-for-dates babies
- On long term steroids or antipsychotics
- Polycystic Ovarian Syndrome or other features of insulin resistance
- Increased BMI ≥ 30kg/m2 (or≥27kg/m2 in Indo-Asian)
- Increased Waist Circumference: ≥94cm (Men) and ≥80cm (Women) (or ≥90cm (Men) and ≥80cm (Women) if Indo-Asian)

Note for woman with previous gestational diabetes:

Hba1c is checked by general practice 3 months post-delivery and at least annually thereafter depending on result (sooner if HbA1c closer to 49).

Interpretation of screening results (adults) https://t2dm.nzssd.org.nz/

HbA1c Result	Fasting Glucose	Non fasting glucose	Diagnosis	Comments
≥50mmol/mol,	≥7.0mmol/L,	≥11mmol/L	Diabetes	Start treatment.
with symptoms	with symptoms		2.00000	Start treatment
				2 Abnormal tests (HbA1c or
≥50mmol/mol, no	≥7.0mmol/L, no		Diabetes	fasting) are required to confirm
symptoms	symptoms		Diabetes	diagnosis, either on the same day
				or subsequent day.
				Offer lifestyle advice. Perform
				CVD risk assessment and follow
41-49mmol/mol	6.1-6.9mmol/L		Pre-Diabetes	guidelines for treatment of risk.
				Repeat checking of HbA1c every
				6-12 months
				Normal range. Repeat HbA1c at
≤40mmol/mol	≤6.0mmol/L		Diabetes Unlikely	next CVD assessment or when
				clinically indicated.

- Use of fasting plasma glucose is recommended when HbA1c results are borderline or further investigation of the result is necessary, such as when a patient has two differing HbA1c results.
- In pregnancy, HbA1c should be included in **first antenatal bloods** but is not useful in the 2nd or 3rd trimester for diagnosis of GDM.
- Fasting plasma glucose remains a valid test for diagnosing people with Type 2 Diabetes,
 including when HbA1c is not appropriate or cannot be used or may be inaccurate in patients
 with haemoglobinopathies with the direction of change depending on the specific diagnosis, as
 this could cause HbA1c to be artificially high or low.

New Zealand is an outlier as in the rest of the world the cut off for diagnosing diabetes is an HbA1c of ≥48mmol/mol, and pre-diabetes is considered to be an HbA1c of 42-47mmol/mol. It is anticipated that the criteria in New Zealand will change to come into line with the rest of the world in the near future.

Screening for diabetes - Children

Australasian Paediatric Endocrine Group recommend that targeted screening should be considered in children and adolescents aged > 10years old or at onset of puberty, which ever occurs first in those who are overweight (BMI \geq 85 and < 95%) or obese (BMI \geq 95%) and who have one or more of the following risk factors:

- maternal history of diabetes or gestational diabetes mellitus during the child's gestation;
- family history of type 2 diabetes in first degree relative;
- race and ethnicity (Māori, Pasifika, South Asian, South East Asian, Middle Eastern, North African and Latino);
- signs of insulin resistance (acanthosis nigricans);
- conditions associated with obesity and metabolic syndrome (hypertension, dyslipidaemia, fatty liver disease, polycystic ovary syndrome, or small for gestational age); and
- use of psychotropic medications

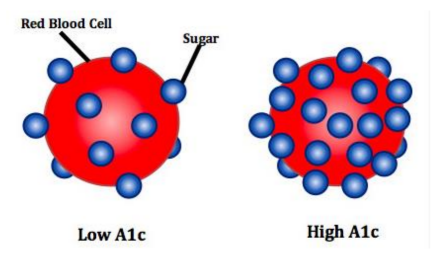
Interpretation of screening results (children)

Diagnosis (glucose measurements)	 Classic symptoms of diabetes or hyperglycaemic crisis and random plasma glucose ≥ 11.1 mmol/L, or Fasting plasma glucose ≥ 7.0 mmol/L, or 2-hour plasma glucose ≥ 11.1 mmol during an OGTT, or HbA_{1c} ≥ 48 mmol/mol (≥ 6.5%)
Other investigations at diagnosis	 Consider diabetes autoantibody testing in all paediatric patients with the clinical phenotype of type 2 diabetes due to higher prevalence of type 1 diabetes — GAD and IA2 are the most commonly available. Insulin antibodies and zinc transporter 8 antibodies can also be measured if negative GAD and IA2, and type 1 diabetes is suspected Insulin and C-peptide measurements are not recommended as glucotoxicity and lipotoxicity can acutely affect insulin secretion

For more guidance, phone 06 753 6139 and ask for the on call paediatric registrar or contact the paediatric diabetes CNS on 027 5577 110

HbA1c

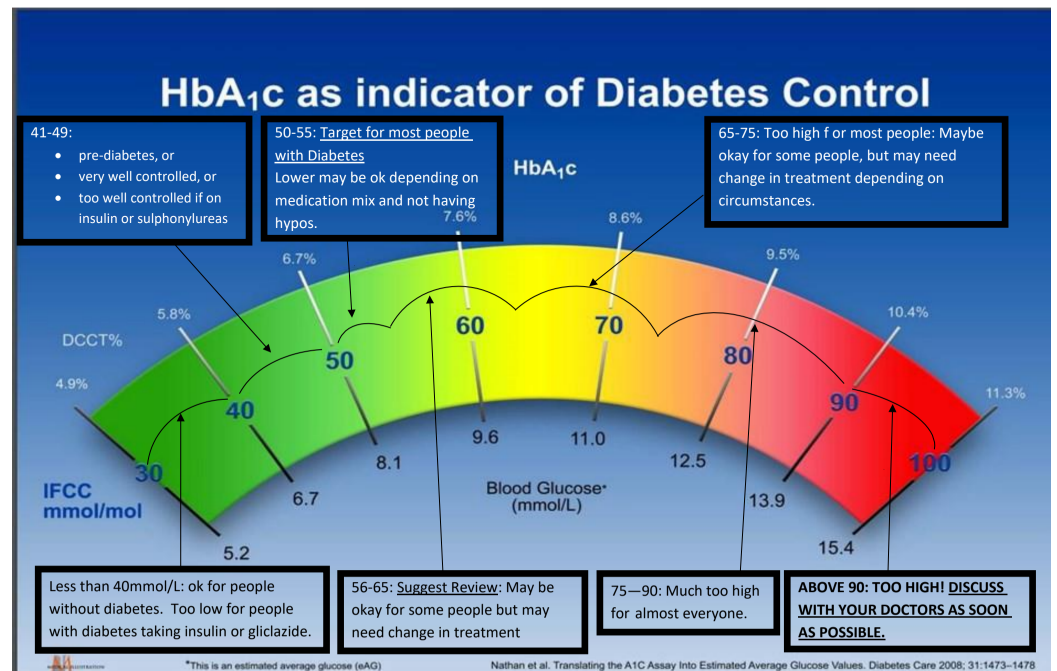
HbA1c is the measure of how much glucose has become stuck to the haemoglobin molecules on the red blood cells. By looking at the oldest red blood cells (remember that red blood cells only live for about 12 weeks) it gives an average of how much glucose has been circulating in the blood for the previous 3 months. This is the preferred test for diagnosing diabetes, and for on going monitoring of control.



For most people in New Zealand, the target HbA1c is \leq 53mmol/mol, however the frail, elderly or those with limited life expectancy may be safer with an HbA1c at 60-65mmol/mol.

Until your patient has an HbA1c to target, it should be monitored every 3 months, and if still above target an escalation of therapy is required to implement change.

Factors which can increase HbA1c	Factors which can decrease HbA1c
Alcohol intake	Erythropoietin, iron or B12 administration
Iron or B12 deficiency	 High ingestion of antioxidants (vits C&E)
Hyperbilirubinaemia	Very high triglyceride levels
Opiate use	Chronic aspirin use
 Splenectomy 	Rheumatoid arthritis
Renal failure	Use of antiretrovirals
	Post blood transfusion
	Anaemia
	Renal failure



*This is an estimated average glucose (eAG)

Nathan et al. Translating the A1C Assay Into Estimated Average Glucose Values. Diabetes Care 2008; 31:1473-1478 Christchurch Diabetes Centre 2009

Clarification of diagnosing and coding

Advice for those people who have an HbA1c that has returned to within normal range.

Once a person is diagnosed with diabetes, **HbA1c** is an indication of <u>control only</u> - not an indicator for change in diagnosis. This is why diagnosis in the first instance needs to be clear and well documented.

- Target HbA1c for most New Zealanders is <53mmol/mol, and with new medications in the toolbox that do not cause hypoglycaemia, it is safe and even ideal to have an HbA1c in the 40s.
- There are a few patients who fulfil the criteria for a diagnosis of diabetes, and having been coded as such, may achieve long-term normalisation of glycaemic control without medication, through lifestyle measures or bariatric surgery.
- In the UK, GPs are advised <u>not</u> to remove such patients from relevant screening registers (including retinal screening) as current evidence shows that all people with diabetes should be screened for diabetic retinopathy for life once there has been a definite diagnosis, excluding gestational diabetes.
- Studies show up to 40% of people who go into remission (using ADA criteria) following bariatric surgery will have relapsed within 5 years.

As yet, there is no criteria for remission of diabetes agreed on in New Zealand. However, in academia,

REVERSAL of diabetes is considered when HbA1c <48mmol/mol with metformin only.

REMISSION of diabetes is considered when HbA1c has been <48mmol/mol for over 12 months with no pharmacological input at all.

If you *do* consider removing the diagnosis of diabetes:

- Clear documentation in the notes by the patients GP as to why the diagnosis was changed
- How the patient is informed of the changes and a record of this occurring
- Consideration of medico-legal issues relating to removing a diagnosis of diabetes if the
 patient fulfilled criteria for that diagnosis at one time, particularly as it relates to past
 medical history for the purpose of medical insurance or life insurance.

HbA1c 41-49

- This can indicate prediabetes.
- If your patient has been previously diagnosed with diabetes, it indicates very good control.

 Someone taking metformin, vildagliptin and empagliflozin or dulaglutide might have an HbA1c in this range and have no risk of hypoglycaemia and requires no down titration of their medication.
- If your patient is prescribed insulin or a sulphonylurea, an HbA1c in this range indicates they have hypoglycaemia reasonably regularly, and their medication should be down titrated.

Type 1 or type 2 diabetes?

Anti-IA2, anti-GAD or anti-ZnT8 can help diagnosis.

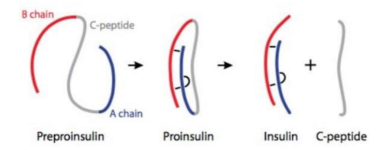
C peptide must always be done with a corresponding glucose result.

If initially managed on diet and oral medications— but are now on insulin,

they have Type 2 diabetes on insulin.

Some background information

C peptide is a part of proinsulin. When the insulin splits off, C peptide is left. It is more stable in the blood stream and therefore a more useful measure than insulin.



In a pancreas that is functioning normally the C peptide level will correspond with the amount of glucose in the blood stream. The more glucose that is present the more c peptide is present.

In type 1 diabetes C peptide can be very low despite a high glucose level i.e. <250pmol/L fasting or <600pmol/L post meal if glucose >8mmo/L.

With type 2 diabetes there are a number of options. If the glucose level is low and C peptide is high this indicates insulin resistance. If the glucose is high, and the C peptide is lower than expected, this indicates insulin deficiency. This is why it is critical to do both tests, C peptide and glucose at the same.

Note that there are other indicators for c peptide checking including insulinoma (may be indicated by a high C peptide and low blood glucose).

GAD stands for Glutamic acid decarboxylase. While this usually acts to make neurotransmitters, it can at times work as an autoantigen. It triggers the autoimmune system to produce autoantibodies to destroy its own beta cells. GAD antibodies can be found in 70 - 80% of people diagnosed with Type 1 diabetes. If at the same time there are autoantibodies to Islet cells (tyrosine phosphatase—related islet antigen 2 = (IA2), it strengthens the diagnosis. Type 1 diabetes is an autoimmune condition.

If you are concerned that the GAD and IA2 have missed the type 1 diagnosis, request a zinc transporter check as well. These are done at Taranaki hospital and in some areas are added automatically if GAD/1A2 come back negative. Request <u>ZnT8 transporters</u>.

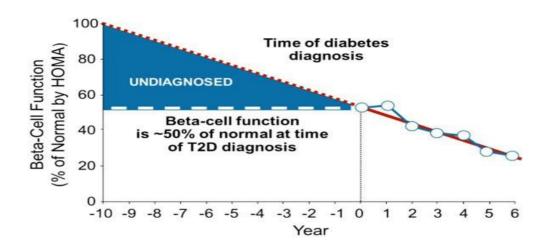
There are more than simply the 2 types of diabetes—MODY, GDM and LADA are a few of the more common ones. If you need more information on these conditions, try this article: CMAJ 2014 Approach to the patient with atypical diabetes; Jun 10; 186(9): 678–684.

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4049991/

PRE-DIABETES

- Diabetes is one of New Zealand's fastest growing long-term conditions
- Annually, 5-10% of individuals with pre-diabetes will progress to T2DM
- Prevalence has been increasing approximately 7% per year for the past 8 years
- The appropriate term to classify patients with HbA1c 41-49 is debated. In 2006, the WHO recommended use of the term "Intermediate Hyperglycaemia" as opposed to "prediabetes", as not all individuals with an HbA1c between 41-49 would proceed to Type 2 Diabetes. (From MOH pre-diabetes and self-management 2016).

Beta cell function can be less than 50% of normal by the time diabetes is diagnosed



Treatment: lifestyle measures are the key treatment Consider: The initiation of Metformin for prevention of diabetes.

ADA consensus 2017 states, 'Metformin therapy prevention of type 2 diabetes should be considered in those with pre-diabetes, especially those with BMI >35kg/m2, those aged >60years, women with prior gestational diabetes mellitus, and/or those with rising A1c despite lifestyle intervention. Long term Metformin may be associated with biochemical vitamin B12 deficiency, and periodic measurements of vitamin B12 levels should be considered in metformin-treated patients, especially in those with anaemia or peripheral neuropathy.

SOME QUESTIONS FOR YOUR PRACTICE:

- Does your practice have a routine protocol for pre-diabetes diagnosis, education, and follow up?
- There are seven possible pre-diabetes classification read codes—encouraging the whole
 practice to maintain one or two of these options simplifies systematic review and
 prompts intervention.
- Recall will depend on results of HbA1c and patient history. If HbA1c close to 49—consider recalling at 3-6 months rather than annually.
- Provide a combination of written and verbal advice.

Some background reading:

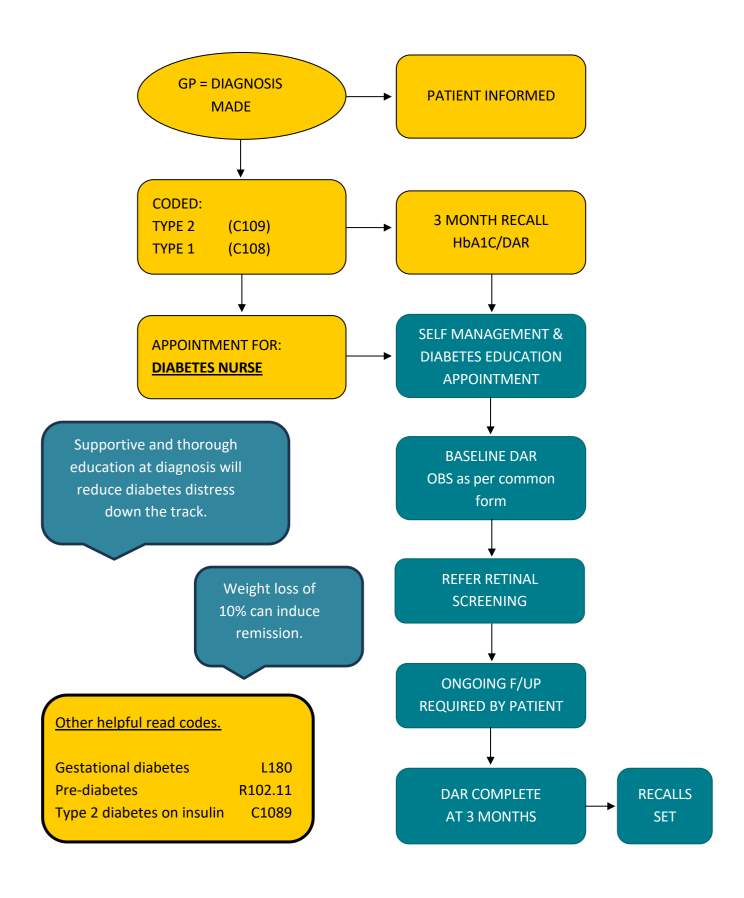
Diabetes Prevention Program (DPP) and the ongoing DPP Outcomes Study (DPPOS) are major studies that changed the way people approach type 2 diabetes prevention worldwide. The DPP showed that people who are at high risk for type 2 diabetes can prevent or delay the disease by losing a modest amount of weight through lifestyle changes (dietary changes and increased physical activity). Taking metformin, a safe and effective generic medicine to treat diabetes, was also found to prevent the disease, though to a lesser degree.

The DPPOS has continued to follow most DPP participants since 2002. To date, the DPPOS has shown that participants who took part in the DPP Lifestyle Change Program or are taking metformin continue to prevent or delay type 2 diabetes for at least 15 years. The DPPOS has also shown that the DPP Lifestyle Change Program is cost effective (costs are justified by the benefits of diabetes prevention, improved health, and fewer health care costs) and metformin is cost-saving (leads to a small savings in health care costs) after 10-years.

Direct and Preview studies have investigated weight loss early in the diabetes journey and its positive impact on blood glucose control, and even preventing diabetes in those with prediabetes.

- Pre-diabetes is not a clinical condition in its own right but is a high-risk factor that can contribute to both diabetes and cardiovascular disease.
- Prevalence of pre-diabetes is particularly high in some population groups for example: Over 40% of people living in the Auckland metro region who are Māori, Pacific, or Indian ethnicity have been identified with pre- diabetes at 35-39 years, and over 50% at 45-49 years (Chan 2015).

INITIAL EDUCATION AND DAR



Patient initial education

How we educate people is as important as what we teach



Assessment: What do they already know?

Establish what the person/whānau already know about Diabetes, build on this when answering questions they may have.



Make it personal for the person with diabetes

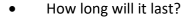
Make information relevant to **their** diabetes, so it makes sense to them—consider their health beliefs and their previous experience/knowledge of diabetes



Consider questions that the person/whānau may not voice

- What caused their Diabetes?
- What is actually wrong?





- How good is my treatment at controlling my Diabetes?
- What can I do to look after myself?

These are just some of the questions that a person can have but be too shy to ask.



Documentation

Document what information you have covered, and any gaps that you believe will need to be discussed at a later time.

Use a keyword to make documentation easier and more consistent. E.g. diabetes education:

- Food
- Checking
- Activity





Goal setting

Small achievable goals are the key to lifestyle change



SPECIFIC

- Define the goal with clear, understandable language
- WHO is involved? WHAT do I want to accomplish? WHERE



MEASURABLE

- Can you track the process and measure outcomes?
- How much, how many, how will I know when at my goal?



ATTAINABLE/ACHIEVABLE

- Is the goal reasonable enough to be accomplished? How so?
- Is the goal out of reach or below standard performance?



RELEVANT

- Is the goal worthwhile and will it meet your needs?
- Is each goal consistent with other goals you have established?



TIMELY

 Your objective should include a time limit. E.g. I will complete this step by month/day/year

Work together to ensure best results.

Person with diabetes Health professional

People are capable partners in their health care

Diabetes Annual Review

- Type 2 diabetes is a progressive condition
- Often treatments are layered in over the years; this means that on-going control is reviewed regularly.
- An annual review is standard care for all people with diabetes.
- An annual review is the screening phase. Then triage—who needs to be seen again urgently, soon, next year? Who needs to be referred to a secondary service such as podiatry, renal or dietetics?
- Treatment is individualized

DAR bloods can be **preset** on MedTech request form, and the pathlab e-request form (right click)

Minimum: HbA1c, Full (non-fasting) lipid profile, Renal function, full blood count urine microalbuminuria, consider B12 if on long term Metformin, and LFT if on Vildagliptin.

Annual reviews are about the person who has diabetes. It is their opportunity once a year to check in and make sure their diabetes management and education is up to date. If we are prepared prior to them arriving, it means that we can focus on what is most important to the person.

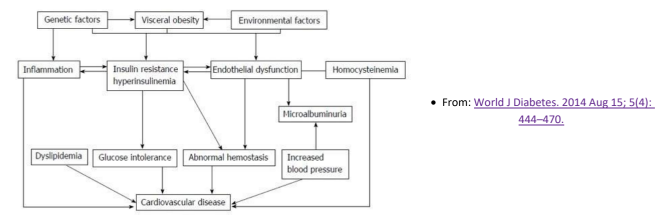
On the next page is a suggested format for preparing yourself for an annual review. Take a photocopy and fill in everything you can prior to the person arriving, and then you will have identified the clinical issues and are free to focus on them. It is most helpful to create a plan forward together.

Example of Diabetes Annual Review on next page:

Diabetes Annual Review Preparation Sheet Patient Name/NHI: Today's Date: **Duration:** Type of diabetes: On-going plan: Comment: Relevant medications: Date most recent results: ひ⇔ひ Hba1c Metformin **Cholesterol Total:** Ratio: Vildagliptin LDL: HDL: Empagliflozin Trig: Kidneys EFGR: Dulaglutide Microalbuminuria: Sulphonylurea *Insulin (type/device/amount) Last result Today's result BP: Weight: Height: **Diabetes Distress Score** WC: DSS2 CVRA: PQH2 Smoking: Exercise: Type duration frequency Food – any changes and goals: Assessment of feet complete and able to self-care: Checking when, recording, changes required: Hypos: YES/NO What used to treat hypos? \bigcirc discuss contraception or pregnancy planning discuss erectile dysfunction Medical alert Driver licence type if Retinal Vaccines due? Shingles (≥65) /COVID/Flu bracelet?: relevant: screening due: Travel Check sites *If on insulin: Storage Sharp disposal Sick day Insulin dose corect Insulin technique Correct pen and insulin Changing needles

Cardiovascular risk

• All people with type 2 diabetes are at higher risk of cardiac disease. The reason for this is multifaceted and not completely understood.



- CVD risk assessment is now based on New Zealand PREDICT study data beginning assessments 10—
 15 years earlier in some at risk groups.
- The aim is to <u>facilitate informed choices</u>, so patient considers early and lifelong healthy lifestyle choices, and when appropriate other treatments.
- <u>Non-fasting blood lipids</u> are now recommended for all treatment decisions and monitoring as long
 as the triglyceride levels are normal. Fasting bloods are only required if triglycerides are elevated
- Full details can be found here: https://www.health.govt.nz/publication/cardiovascular-disease-risk-assessment-and-management-primary-care
- CVD risk assessment and management is a recommended component of the annual diabetes review, in people with type 2 diabetes.
- The new risk prediction equations for people with type 2 diabetes include: duration of diabetes,
 BMI, eGFR, ACR, HbA1c and hypoglycaemic medications; in addition to the risk factors in equations for people without diabetes.
- No specific risk equations are available for people with type 1 diabetes, although the same main disease variables (diabetes duration, renal disease, glycaemic control) apply as for type 2 diabetes.
 CVD risks for this group are substantially higher than for people with type 2 diabetes (50% higher in men and up to 90% higher in women).
- Encourage a healthy lifestyle (smoking cessation, healthy diet, regular physical activity, optimal weight) in everyone.
- Optimise glycaemic control to an appropriate level in consultation with the individual patient.
 Agreed target range will generally be more stringent in younger and fitter patients (e.g. HbA1c 48-53, FBG 5-7mmol/L without hypos) than older, co-morbid or frail patients and those prone to hypoglycaemia (e.g. HbA1c 55-64, FBG 7-9mmol/L without hypos).

Hypertension

A word from the experts!

Between 70 and 80 % of people with diabetes have hypertension.

The coexistence of diabetes and hypertension worsens clinical outcomes with respect to both microvascular and macrovascular disease. Diabetes management should therefore be comprised of a multifaceted approach that targets optimal blood pressure and lipid management in addition to glycaemic control. The pathophysiology of hypertension in diabetes involves maladaptive changes and complex interactions between the autonomic nervous system, mechanical forces, renin-angiotensin-aldosterone system as well as individual and environmental factors. Multiple high-quality randomized controlled trials have shown reduced morbidity with reduction of elevated blood pressure in people with diabetes.

What this means in practice is — please treat hypertension!

Treatment needs to be individualised but <130/80 is the recommendation for most people, although <125/75 likely to benefit younger patients.

Causes of hypertension

Sedentary lifestyle, increased calories and insulin resistance:

 Increased adiposity = increased insulin resistance = oxidative stress and inflammation = vascular stiffness = hypertension

Increased intra vascular volume:

 Mismatch between sodium intake and sodium loss = increased sodium concentration = increased venous return = elevated arterial pressure

Premature vascular aging:

 Structural changes to the lumen of small vessels = vascular remodelling, inflammation, peripheral vascular resistance, decreased elasticity = hypertension

Autonomic nervous system deregulation:

• Increased sympathetic activity = increased heart rate, ventricular contraction, peripheral resistance, and fluid retention.

Role of innate and adaptive immunity:

- Involved in angiotensin II and aldosterone induced hypertension. Renin Angiotensin Aldosterone system
- Angiotensin II is a potent vasoconstrictor and has a direct effect on the vascular smooth muscle also promotes sodium and water retention

Dyslipidaemia

The New Zealand Heart foundation has a great explanation of Cholesterol.

Cholesterol (Matu Ngako) is a type of fat that circulates in your blood. Your body needs some cholesterol to work properly, but when you have high cholesterol (hyperlipidaemia) it speeds up the process of atherosclerosis.

This is when plaque builds up in your artery walls, narrowing arteries and restricting blood flow. Eventually the plaque can get so large and hard that it cracks, and a clot forms over the plaque. If this clot blocks an artery, it can cause a heart attack or stroke.

There is only one type of cholesterol, but it is carried around the body by different 'carriers' (lipoproteins) that have different tasks. There are two types of carriers: LDL (low density lipoprotein) and HDL (high density lipoprotein).

LDL-cholesterol L-L-L (LDL is Lousy and it needs to be Low)

- The role of LDL in the body is to transport cholesterol to all organs for use in building cells.
- LDL is like a large 'dump truck', and dumps cholesterol in the artery walls.
- Known as 'bad' or 'lousy' cholesterol.

For high-risk individuals an LDL-C target of 1.4mmol/L or lower is recommended.

For intermediate-risk individuals the benefits and harms of lipid-lowering drugs should be presented and discussed to allow an individualised informed decision about whether to start treatment. A target LDL-C reduction of 40% or greater is recommended if drug treatment is commenced.

HDL-cholesterol H-H-H (HDL is Happy and it needs to be High)

- HDL cholesterol is like a 'pick up' truck that picks up and transports cholesterol from the blood stream to the liver, which gets rid of it.
- The role of HDL in the body is to carry extra cholesterol away from the arteries to the liver.
- Known as 'good' (or happy) cholesterol.

Triglycerides are the most common form of fat in your body. The role of triglycerides is to store and transport fat in the blood. Extra energy from food and alcohol that your body does not need is changed to triglycerides. High triglycerides increase your risk of heart attack and stroke.



Total cholesterol is a rough measure of all the cholesterol and triglycerides in your blood.

Total cholesterol/HDL cholesterol ratio is the ratio of your total cholesterol to your HDL cholesterol. This ratio is used to measure your risk of heart attack and stroke..

Diabetes recall process

- What system does your practice have in place?
- Is it one person or multiple people responsible for recalls?
- People with type 1 diabetes may or may not want an annual review at the GP—offer them a choice and aim to have a DAR in primary health 6 months after the specialist review.

DAR

- First recall letter and blood test form sent
- Enclose blood test form

CONTACT

- Phone calls, emails, text and portal contacts
- Document each contact attempt
- You could set up a key word for this

ONGOING

- Task GP to inform outstanding DAR
- Aim for opportunistic DAR or continue with contact attempts



- This is MOST important—what happens after the DAR?
- How soon do you need to phone or see the person to assess goals, HbA1c, BP or any other issues that were raised in DAR. Who follows this up?

LIFESTYLE

FOOD ACTIVITY SLEEP STRESS

Numerous aspects of lifestyle and wellness affect a person's diabetes. There are real constraints in general practice as to how we can support the patient to live as well as they possibly can.

There are some key concepts that support our understanding of diabetes management. Each of you will find different ways of explaining these concepts to patients. Remember cornerstone of treatment for type 2 diabetes is:

healthy food choices, physical activity, medication



There is no conclusive evidence to suggest one dietary strategy is more effective than any other for achieving sustained weight loss and improvements in glycaemic control. The choice of dietary strategy will depend on many factors but particularly patient preference, tolerance, nutritional needs, income, comorbidities and cultural suitability.

Some recommended sites for lifestyle information

https://t2dm.nzssd.org.nz/Section-88-Healthy-eating (has links to some dietitian resources for practice nurses)

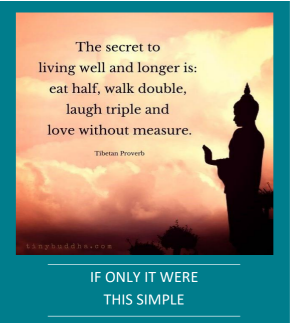
https://www.diabetes.org.nz/





Eating healthy food is an important part of self-managing diabetes

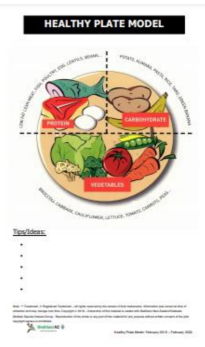




Nutrition resources

There are so many resources to choose from—be selective. Aim for no more than 5 resources you use a

lot. Here are a couple more favourites.





Healthy Food Guide has a portion size poster as well—it has more detail than this one from the heart foundation

Try this at







Remember that people get confused if they are told different health messages. Keep to evidence-based health/food messages. Ensure as a team you are all saying the same things about food. Fad diets typically

promise quick weight loss, restrict a specific food group, require expensive

supplements, and are short lived.

If it sounds too good to be true—it probably is!

DIETARY INTAKE

All people with diabetes need to know that carbohydrates impact on their BGL.

Make an opportunity to assess a person's food intake and prior knowledge level.

Carbohydrate (CHO) foods provide not only "carbs", "energy", and "fuel" but also many health promoting nutrients from fibres to phytonutrients. Discuss whole verses refined for their health benefits.

Discuss some common labelling misconceptions. (No added sugar, organic, fat free).

Make sure that patients are aware that increased calorie consumption over their individual requirements will increase their weight.

Enjoy a variety of nutritious foods every day including:

- plenty of vegetables and whole fruit, wholegrain foods, and those naturally high in fibre
- some milk and milk products, mostly low and reduced fat
- some legumes, nuts, seeds, fish and other seafood, eggs, poultry (e.g. chicken) and/or red
 meat with the fat removed. Consider swapping some animal protein for plant-based protein
 meals.
- Choose and/or prepare foods and drinks with unsaturated fats instead of saturated fats, foods that are low in salt (sodium); and if using salt, choose iodised salt, foods with little or no added sugar, and foods that are mostly 'whole' and less processed.
- Make plain water your first choice over other drinks minimise sugar and alcohol containing drinks.
- No alcohol is good for you, it increases your risk of atrial fibrillation and stroke, and may
 increase your risk of heart failure and IHD. If you choose to drink alcohol, keep your intake
 low. Stop drinking alcohol if you could be pregnant, are pregnant or are trying to get
 pregnant.
- Buy or gather, prepare, cook, and store food in ways that keep it safe to eat.
 - To the best of your ability avoid processed foods.

An Example: if a person's favourite food is pizza and they eat this 3 times a week, could they eat it once a week, or eat a smaller portion (and add a salad on the side), or change the choice they make from Hawaiian stuffed crust x large (1512 KJ) to large slice thin and crispy (626 KJ).

Basal Bolus insulin does work best with great CHO knowledge and when teaching our patients we should focus on timing, type and amount as this is very relevant to achieving control and balance in their diet.

Some people require more in-depth information e.g. education on Glycaemic Index, being able to read nutrition labels, in general how much CHO they consume at each meal. Dietitians specialise in teaching CHO management whilst maintaining good nutrition, and patients on or starting insulin meet criteria for referral. People prescribed Empagliflozin have a minimum CHO requirement of 130g per day.

For further support patients can be referred to the Pinnacle MHN Extended Care Team Community
Dietitian via BPAC.



Basic CHO Sheet

Bread Bread	Carbohydrate	Vegetables	Carbohydrate
1 slice – check the label	10 – 20 g	1 med potato (135 g)	15 g
1 medium slice Molenburg	13 g	1 cup mashed potato	30 g
1 toast slice Molenburg	17 g	1 small piece kumara (135 g)	18 g
1 slice Vogel	18 g	1 whole cob corn	30 g
1 med slice white bread	13 g	½ cup creamed corn	23 g
1 toast slice white bread	17 g	½ of 420g can baked beans	23 g
1 slice your usual bread	g	1 cup cooked dried beans	30 g
1 bread roll long/hamburger	30 g	1 parsnip (22 cm long 160 g)	20 g
1 bread roll med bakery	20 g	Yams (135g)	17 g
1 crumpet	20 g	1 cup thick vegetable soup	20 g
1 large pita bread	40 g	Taro (135 g)	33 g
1 wrap – check the label	15-40 g	Plantain/Green banana (135 g)	33 g
1 English muffin	30 g		
Cereals		Fruit	
1 Weetbix	10 g	1 banana	30 g
1 cup cooked porridge	30 g	2 raw apricots/plums	10 g
½ cup raw rolled oats	25 g	150 gm berries	10 g
1 cup muesli - check the label	90 g	½ cup stewed fruit (in juice)	10 g
1 cup just right/ light n tasty	54 g	1 cup grapes/cherries	25 g
Weetbix Bites (45 g)	32 g	1 apple	15 g
1 cup All Bran flakes	28 g	1 orange	10 g
1 cup Kornies/rice bubbles	25 g	2 medium feijoa	5 g
1 cup Nutrigrain	25 g	1 slice fresh pineapple	13 g
Pasta/Rice		Dried fruit	
1 cup cooked pasta	40 g	4 dried apricot halves	10 g
1 cup cooked rice	50 g	1 cup dried fruit	100 g
1 pkt instant noodles	50 g	1 tablespoon raisins/sultanas	10 g
½ of 420g can spaghetti	23 g	4 dates	20 g
Dairy products		Crackers biscuits baking	
1 glass milk (200 ml)	10 g	1 med cracker (check the label)	5 g
1 pottle diet yoghurt	10 g	1 large cracker	10 g
1 pottle regular yoghurt	25 g	1 plain biscuit (arrowroot, round wine)	5 g
1 ice-cream slice	10 g	1 large biscuit (digestive)	10 g
1 glass soy milk (200ml)	10g	1 rice wafer	10 g
Extras		1 sweet muffin (small 80 g)	40 g
1 pie	45 g	1 scone (small 80 g)	30 g
1 cup macaroni cheese	35 g	1 pikelet (small 25 g)	10 g
1 hamburger – regular	30 g	1 cup flour	90 g
¼ pizza (2 slices)	60 g	1 pkt crisps (small 40 g)	18 g
2 Sushi rolls	20 g	1 pkt crisps (large 150 g)	68 g
1 serve hot chips - med	60 g	1 muesli bar (read the label)	15 – 40 g

Food/glucose activity record sheet

Please record as much information as you can the 4 days before your appointment. You do not need to change your normal pattern of eating, but please write down what you do eat. It also helps to record your blood glucose levels before the meals and 2 hours after the meals. If you normally snack between meals, eat AFTER you have done the 2-hour glucose check.

Remember - there is no such thing as a bad level, as all the information can be used to help you manage your diabetes better.

	Day one	Day two
Before meal blood glucose level		
Breakfast		
2 hours after meal blood glucose level		
Morning tea		
Before meal blood glucose level		
Lunch		
2 hours after meal blood glucose level		
Afternoon tea		
Before meal blood glucose level		
Dinner		
2 hours after meal blood glucose level		
Supper before bed		

	Day three	Day four
Before meal blood glucose level		
Breakfast		
2 hours after meal blood glucose level		
Morning tea		
Before meal blood glucose level		
Lunch		
2 hours after meal blood glucose level		
Afternoon tea		
Before meal blood glucose level		
Dinner		
2 hours after meal blood glucose level		
Supper before bed		

Bring these records to your appointment Remember to bring your glasses if you wear them!

Physical activity

Physical activity is a key component in diabetes management It increases insulin sensitivity and increases consumption of glucose by the cells.

The latest NZ MoH guidelines are for ADULTS:

Do at least 2 ½ hours of moderate or 1 ¼ hours of vigorous physical activity spread throughout the week.

Moderate intensity activity causes a slight, but noticeable, increase in breath and heart rate. You can still carry on a conversation.

Vigorous intensity makes you out of breath – you can't do these activities and chat at the same time.

For extra health benefits, aim for 5 hours of moderate or 2 ½ hours of vigorous physical activity spread throughout the week.

Do some muscle-strengthening activities on at least 2 days each week. More information can be found here: https://www.health.govt.nz/our-work/preventative-health-wellness/physical-activity

GREEN PRESCRIPTION – discuss a referral to green prescription as a first step. Sport Taranaki provide this service and patients can be referred via BPAC

Exercise may impact on the BGL for up to 24 hours. People need to be aware of the potential of a hypo following moderate or vigorous exercise.

Ideally adults with Type 2 diabetes should do both aerobic and resistance exercise training for optimal glycaemic and health outcomes (ADA,2016).

Pre exercise health check by GP is advisable for people with diabetes if previously sedentary or morbidities present.

There are many opportunities for people to increase their activity.

It can be as simple as parking further away from work, walking upstairs not taking the elevator, sit and

be fit activities.

Dancing, walking, swimming, waka alma, in2hula, marathons, TriMāori, hiking... the list is endless. Be aware of what opportunities are in your local area.

It must be affordable, achievable, and often. Doing a small amount of activity regularly is better than a huge effort once in a while.

Consider setting a goal, invite a friend, gradually increase the intensity.

Some people are motivated by technology—pedometers are cheap, fitness watches and apps such as my fitness pal or charity miles.

Reduces risk of developing diabetes complications

Regular Physical Activity

Controls weight

Lowers blood glucose levels pressure and cholesterol levels

Encourage people to start any new exercise slowly and aim to increase.

Sleep

Reduced sleep is now widespread, usually as an accommodation to a busy lifestyle and the use of technology intruding into sleep time.

The Sleep Heart Health Studied those with a self-reported short sleep duration (<5 hours) had an increased risk of type 2 diabetes compared to those with a sleep duration of 7—8 hours.

You can find more information about sleep hygiene on the Healthify NZ page:

https://healthify.nz/health-a-z/s/sleep-problems/

Rate 1...... 10 how good was you



Sleep tips tracker



Having difficulty sleeping? There are lots of things you can do to help get a better night's sleep. Use this form to record the changes you are making – remember the more changes you make, the more likely you are to get a good night's sleep. For more copies, visit www.hn.org.nz/takecharge

If blood glucose levels are high overnight, nocturia can be an issue and further interrupt sleep patterns. Restless leg syndrome is sometimes experienced by people with diabetes as is cramping and neuropathic pain. All of these issues will improve with improved blood glucose control.

Consider assessment for sleep apnoea

Obstructive sleep apnoea (OSA) alters glucose metabolism, promotes insulin resistance, β cell dysfunction, and is associated with development of type 2 diabetes. This appears to be independent of obesity. Chronic exposure to intermittent hypoxia and other pathophysiological effects of OSA affect glucose metabolism directly and treatment of OSA using a CPAP machine can improve glucose homeostasis. Lifestyle interventions has a significant impact on Apnoea hypopnoea index (AHI) in people with type 2 diabetes.

If people are struggling with sleeping well, encourage them to talk to their GP.

Stress

KEY POINTS for the person with diabetes

- Stress is a part of our daily lives
- Stress becomes unhealthy when it begins to make us less able to manage our physical or psychological health, or other factors in our lives such as our work and relationships
- Stress can be caused by physical factors (like an injury or illness) or psychological or social factors (unresolved work issues, bereavements, moving house, unresolved relationship problems)
- For many people with diabetes, stress can cause blood glucose levels to rise. Learning strategies to deal with stress may lessen this effect
- Having diabetes is in itself a major source of stress. People with diabetes have higher rates
 of anxiety and depression. Learning how to manage stress and treating these skills as a
 priority, can help a person cope with stress more effectively
- There are practical things that can be done to reduce stress, such as learning relaxation techniques, learning different ways to respond to stress, identifying situations that cause stress and choosing to avoid them, and making changes to your life that increase your enjoyment level
- Developing a positive coping style may help you deal more effectively with stress

Stay vigilant for diabetes burnout and distress.

Screening for psychological problems and diabetes burn out has not been happening systematically. In order to address this, the **PHQ2** Depression questionnaire and **DDS2** are now a routine aspect of the diabetes annual review. These give more focus to the persons feeling of coping with the rigorous demands of living with diabetes. If the person you are seeing scores highly when answering these questions, your role is to support them to achieve healthy habits that benefit their physical body as well as their mental health – good food, good exercise, good sleep habits, meditation and referral to the GP highlighting your concerns

Adult weight management

Referrals can be made to the dietitian within the Pinnacle MHN extended care team via BPAC.

Prior to the referral, please assess that the person is at a stage where they want to address their diet and lifestyle.

Encourage realistic weight loss targets of 5-10% body weight.

All of the lifestyle topics addressed here are great for the whole population.

Be the champion for healthy lifestyle messages.

MICROVASCULAR COMPLICATIONS

It is essential to identify diabetes related complications by early and regular screening and this includes gum disease, foot examination for neuropathy and ulceration, kidney damage and eye disease. Patient education is necessary in all of these areas to avoid impaired quality and length of life and burden of costs to secondary health care services. (MOH, 2015. Living well with Diabetes).

While the <u>UKPDS</u> is an older study in today's terms, it still a landmark for reducing complications. The key messages are:

- retinopathy, nephropathy, and possibly neuropathy are benefited by lowering blood glucose levels in type 2 diabetes with intensive therapy. The overall microvascular complication rate was decreased by 25%.
- Tight control of blood pressure reduced the risk of any non-fatal or fatal diabetic complications and of death related to diabetes

This is why there is Think about how a target for good to communicate glycaemic control, this to your which is HbA1c 53 patient for most people. Macrovascular disease Microvascular disease Transient ischaemic attack Diabetic retinopathy Stroke non-proliferative proliferative macular oedema Angina Myocardial infarction Cardiac failure Microalbuminuria Macroalbuminuria End-stage renal disease Erectile dysfunction 11 Autonomic neuropathy

Peripheral

vascular

disease

Peripheral neuropathy

Osteomyelitis

Amputation

Renal disease

All people with type 2 diabetes should have eGFR and Albumin/Creatinine ratio assessed annually and more often if abnormal (3 -6monthly)

If the patient has albuminuria, then this result should be repeated one or two times over the next 3 months to confirm the result. Exclude causes such as UTI, severe hyperglycaemia, heart failure, vigorous physical activity, contamination with blood, or other kidney disease i.e., concomitant haematuria is present. Confirm with another test.

The combination of a low eGFR and albuminuria/proteinuria means that the patient is at greater risk of developing end-stage renal failure, compared with patients who have a low eGFR, but no albuminuria or proteinuria.

				Persistent Albuminuria Categories Description and Range		
				A1	A2	А3
				Normal to mildly increased	Moderately increased	Severely increased
				<30 mg/g <3 mg/mmoL	30–300 mg/g 3–30 mg/mmoL	>300 mg/g >30 mg/mmoL
	G1	Normal or high	≥90			
73 m²)	G2	Mildly decreased	60–89			
nL/min/1	G3a	Mildly to moderately decreased	45–59			
GFR Categories (mL/min/1.73 m²) Description and Range	G3b	Moderately to severely decreased	30–44			
GFR Ca De	G4	Severely decreased	15–29			
	G5	Kidney failure	<15			

Kidney Disease: Improving Global Outcomes (KDIGO) 2017 Clinical Practice Guideline classification chart for albuminuria.

For people with diabetes and CKD in the Whatu Ora - Taranaki region there is a clinic run by the hospital renal service. Referral is via BPAC.

PRACTICE POINTS

- People with confirmed microalbuminuria should be treated with an ACE inhibitor or an ARB whether or not hypertension is present.
- Good glycaemic control in patients with CKD and diabetes is essential to prevent or delay the progression of diabetic nephropathy, and to reduce cardiovascular risk.
- Empagliflozin (Jardiance) is the preferred second line therapy and funded for patients with renal complications and eGFR ≥20 mL/min.
- Dulaglutide (Trulicity) would be the next preferred therapy if empagliflozin is not tolerated, and can be used down to eGFR ≥15 mL/min.
- An HbA1c ≤53 mmol/mol is generally a target for patients with CKD and diabetes.
 However, there is an increased risk of hypoglycaemia with insulin or a sulphonylurea for older patients living alone, those with comorbidities or limited life expectancy, and a target HbA1c ≥ 53 mmol may be more appropriate. This should be decided with patients using a shared decision-making approach.
- In patients with advanced stage 4 and stage 5 CKD the risk of hypoglycaemia with insulin or sulphonylurea use is also clinically relevant, and less intensive glycaemic control but with close monitoring is often required.
- KDIGO guidelines now recommend that a Metformin dose of 2g per day is safe when the eGFR is ≥45 mL/min. Metformin should be avoided altogether in patients with an eGFR ≤15 except under the close supervision of a nephrologist.

Using ACE inhibitors

- When renal disease is present, start an ACEi or ARB and maximise dose before starting another agent.
- If an ACEi is commenced, the eGFR and serum potassium should be measured 7—10 days later to ensure there is no further decline in eGFR.
- If reduction in eGFR ≤ 25% and eGFR stabilises within 2 months of initiation, continue the ACEi.
- A reduction > 25% after starting ACEi may indicate renal artery stenosis. Reduce or stop the ACEi and consider nephrology assessment.
- If potassium > 6 mmol/L, stop the ACE inhibitor and switch to an ARB

Some extra notes about treatment of renal disease:

- Any evidence of renal disease based or decreasing eGFR should be treated with urgency.
- Loop diuretics may be used instead of or in combination with thiazide diuretics in patients with significant renal impairment (eGFR < 45 ml/min/1.73m2).
- People with type 2 diabetes are more prone to nephrogenic anaemia (from no other cause)
 and eligible for erythropoietin for eGFR<45. Hb <100

Opinions vary as to when metformin should be reduced in the presence of renal impairment, but all agree that metformin is contraindicated when eGFR<15.

A helpful resource:

'Chronic kidney disease (CKD) management in general practice': https://www.kidneys.co.nz/resources/file/kidneyhealth_complete_pgs-2.pdf

Chronic Kidney Disease (CKD)Management in General Practice

End stage renal disease (ESRD)

Historically this has been the domain of the renal unit, but at times general practice will be asked to manage blood glucose levels. This needs to be done on an individual basis. Some general points are:

HbA1c target is moderate. Expert opinion recommends HbA1c between 53-75mmol/mol. Targeting lower HbA1c levels may exacerbate mortality risk in dialysis patients with underlying illness and malnutrition.

Up to 1/3 of people with ESRD experience 'Burn- out diabetes'. There are a number of factors that contribute to a spontaneous resolution of hyperglycaemia—the HbA1c returns to normal without the use of treatment.

Many glucose lowering drugs and their active metabolites are metabolised and excreted through the kidneys and will therefore require dose adjustment in the dialysis patient.

Limitations of HbA1c in dialysis patients: ESRD related factors can cause falsely high HbA1c values (due to high urea levels and metabolic acidosis) as well as falsely low values (due to anaemia, repeated blood transfusions, haemoglobinopathies, use of erythropoietin-stimulating agents and malnutrition). Despite this, renal guidelines from both the UK and US advise use of HbA1c to monitor glycaemic control in dialysis patients.

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3960718/pdf/nihms558834.pdf

Autonomic neuropathy

This can affect the sympathetic and parasympathetic functions in people with diabetes involving cardiovascular (silent MI), gastrointestinal (gastroparesis, constipation), genitourinary (neurogenic bladder/erectile dysfunction), metabolic (hypoglycaemia unawareness), sudomotor (excessive sweating) and ocular changes (small pupil that constricts poorly to light).

PRACTICE POINT

It is important when screening patients at annual diabetes review you can recognise the symptoms of autonomic neuropathy and refer the patient on for further medical assessment

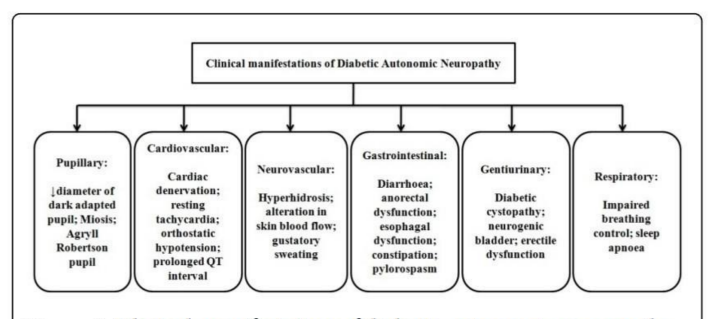


Figure 4: Clinical manifestation of diabetic autonomic neuropathy

Periferal neuropathy

- Diabetic peripheral neuropathy is one of the most common long-term complications of diabetes.
- It develops in up to half of all people with diabetes and is one of the main risk factors contributing to foot ulceration and eventual amputation.
- Basic foot education at an annual review can prevent some issues developing.
- It is the main cause of non-traumatic lower limb amputation, which is a result of a combination of decreased sensation and reduced arterial supply.
- Assessing for peripheral neuropathy is a routine part of ongoing care for patients with diabetes.
- Treatment of diabetic neuropathy includes optimal control of hyperglycaemia, appropriate foot care (often involving input from a podiatrist), and symptomatic management of any neuropathic pain.

A very comprehensive article on Peripheral Neuropathy can be found here: https://bpac.org.nz/BPJ/2014/June/diabetic-peripheral-neuropathy.aspx

For a thorough foot check:

- Take off the both the persons shoes and socks you need to compare feet!!!
- A general inspection of the feet and the patient's footwear
- Musculoskeletal assessment for deformity (including Charcot arthropathy)
- Neurological assessment
- Vascular assessment of the feet, and assessment of the heart rate and blood pressure

General inspection of the feet notes.

Examine both feet and check the condition of the skin, particularly looking for erythematous areas, dryness, flakiness, thickness, cracking, callus formation, infection, and ulceration. Dermatological changes, such as dry or scaly skin may be secondary to a degree of autonomic dysfunction which can begin distally.

There may also be abnormalities of sweating or circulatory instability in the feet, e.g. a hot or cold foot. Heavy callus formation over the pressure points of the foot and signs of localised rubbing or friction, blisters or erythema can also be an indication of inappropriate footwear. Foot ulcers are not caused by neuropathy alone but can occur without injury once hard callus is present over pressure points. If a patient has a loss of sensation in the foot, there will be prolonged and increased forces on the callused areas which then increases the risk of tissue breakdown and ulceration.



Checking for sensation in feet

Monofilament

Monofilament testing uses a 10 g monofilament to assess a patient's ability to feel light pressure at a number of separate sites on the foot. The New Zealand Society for the Study of Diabetes guidelines suggest the examination of 12 sites in total – six on each foot (Figure 1), although some clinicians believe that fewer sites are required, e.g., four sites on each foot. If the patient cannot detect the light pressure at more than one of the designated checking sites, then loss of protective sensation is deemed to be present.

To perform the check the patient is placed supine with bare feet (or their feet raised on a stool in front of the clinician). The use of the filament should be demonstrated to the patient on their upper arm. Ask them to close their eyes and say "yes" when they can feel the filament. The filament should then be placed against the foot, avoiding areas of callus if possible, and pressed until the patient indicates they can feel it, or until the filament bows (Figure 1). The filament should be pressed against the foot slowly over three seconds, not tapped. Site selection should be random and not predictable by the patient.





 $\textbf{Figure 1:} \ Recommended sites for cutaneous sensory pressure perception testing using a monofilament. Monofilament bent to form a C shape.$

N.B. It is recommended that a monofilament is not used on more than ten patients in 24 hours, as they may buckle. The monofilament should also be replaced on a regular basis to ensure it still has a 10 g pressure. In addition, the monofilament should be cleaned with alcohol after each use.

Available to purchase from:

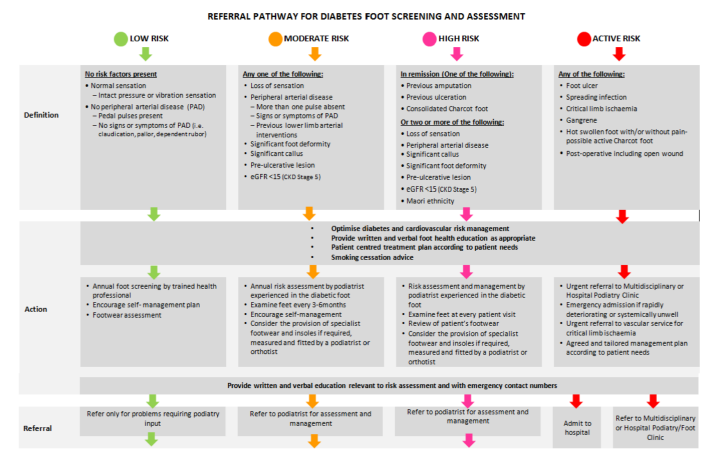
https://www.mediray.co.nz/diabetes/shop/peripheral-neuropathy/neuropen-monofilaments-10g-box-5-gds-nt0104/ or source from an online retailer such as www.aliexpress.com

Ipswich Touch Test

If a monofilament is not available, it is still important to examine and touch the feet. The Ipswich Touch Test involves lightly TOUCHING or RESTING (not brushing, tapping or poking) the tip of the index finger for 1-2 seconds at the same points indicated above. The online DAR form has the Ipswich Touch Test as an option, but defaults to the monofilament test as this is the gold standard assessment. This video shows you how to do the touch test https://www.youtube.com/watch?v=KbMljfRubvQ

Vibratip

The idea with vibration checking is that sensation is the first sensation to be impacted by neuropathy, rather than pressure—so the changes in the foot could be picked up earlier than with a monofilament. Vibratip should only be used in conjunction with a monofilament, not as an alternative.



Updated Referral Pathway 04/04/2017 Adapted from the Foot Action Group (Scottish Diabetes Group) by PodSIG (NZSSD) © 2016

A clearer copy of this diagram can be found here:

https://nzssd.org.nz/content/17_12_2_REFERRAL_PATHWAY_FOR_DIA.pdf

BPACs for active foot clinic can be done if the criteria below are met.

Select the most appropriate option for your patient depending on where they live.

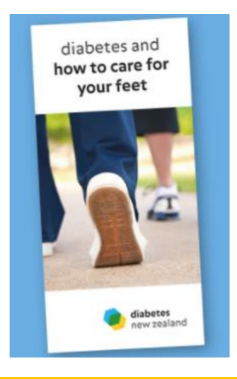
Taranaki DHB Main Referral Centre Podiatry - Secondary (Central/South) Taranaki DHB Main Referral Centre Podiatry - Secondary (North) Diabetes - secondary services (DHB) podiatry referral This service is for diabetic patients with active foot complications and high risk feet. active ulceration spreading infection critical ischaemia gangrene, or; unexplained hot, red, swollen foot with or without pain Include in referral (if available): last diabetic foot check (screening) last three months BPs (screening) last three HbA1c readings BP / height / weight (automatically included if in screening) other relevant clinical history, e.g. consultation For more information please view the Referral pathway for diabetes foot screening and assessment.

Patient education is essential

Encourage your patient to check their feet daily so they will immediately know if there is a change

- After a shower or bath, when putting on shoes and socks, or when getting into bed
- If they can't check their own feet, get a family member to check for them, or they could use a mirror on the floor to check the underside of the foot
- Check between the toes, nails and skin
- Compare the feet colour, temperature, sensation
- Choose shoes that fit have a wide and broad toe, a heal counter, don't rub and are preferably laced or have Velcro.
- Socks are advisable to prevent rubbing (wool or cotton)







You will find some more useful patient resources at the diabetes New Zealand website:

https://www.diabetes.org.nz/

Retinopathy

- Classified into non-proliferative, proliferative, and macular oedema
- Is the leading course of blindness in the developed world countries
- Diabetic retinopathy is a microvascular complication of both type 1 and 2 diabetes
- About 20% of patients with type 2 diabetes have retinopathy at diagnosis.
- The prevalence is strongly related to the duration of diabetes, prolonged hyperglycaemia and the presence of nephropathy, hypertension and dyslipidaemia
- National and international studies show that between 30-40% of people with diabetes have retinopathy
- Severe retinopathy may be present with normal vision
- Educate patients to seek prompt medical assessment if there is any visual deterioration
- Very rapid improvement of glycaemic control can result in short-term worsening of retinopathy. Inform the ophthalmology department if a patient with established retinopathy is being taken from poor control to tight control quickly in case laser treatment becomes indicated
- In the longer term, the risk of retinopathy progression is decreased by tight glycaemic control.
- Refractive errors occur as the lens changes shape with elevated blood glucose

This means that when the glucose levels are high, they may experience blurred vision.
This is reversible. Retinopathy is not reversible.

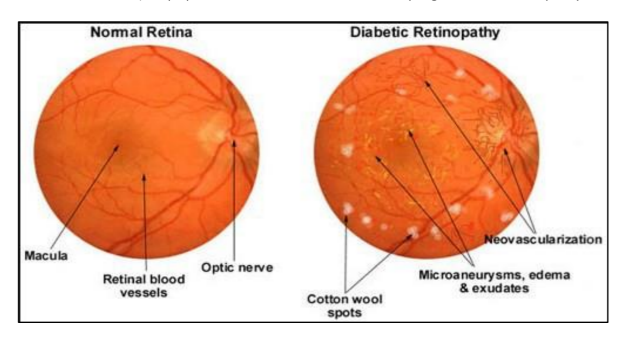
Arrange regular retinal photography:

- For type 2 diabetes, start from diagnosis.
- For type 1 diabetes, start 5 years after diagnosis, not if aged < 12 years.
- If no retinopathy, screen every 2 years— 3 years.
- If retinopathy present, screen at least annually for changes
- If you are not sure when they last had an eye photograph—phone the Taranaki Eye Centre on (06) 758 3553.

ACTIVE MANAGEMENT:

Referral to the eye clinic can be concerning to people; they need good explanations of what is happening and why.

• Active management of all diabetes risk factors (e.g. glycaemic control, blood pressure, lipids, renal function) helps prevent the onset and reduce the progression of retinopathy.



Retinal Photo Screening

Use BPAC for retinal screening:

Entry criteria:

all individuals with a confirmed clinical diagnosis of diabetes

Referral must include:

Most recent labs

HbA1c Creatinine/eGFR /ACR

Type of diabetes

Duration of diagnosis or year of diagnosis

History of foot ulcers

Include BP / height / weight if available (automatically included if in screening).

Note: Gestational diabetes lasts the length of pregnancy – these women will not develop diabetic retinopathy within 9 months and do not meet the service entry criteria.

Normal visual acuity does not preclude sight-threatening retinopathy.

Cataracts are more common in diabetes, seen at a younger age, and progress more rapidly.

Erectile dysfunction

Erectile dysfunction is defined as the inability to get and maintain an erection

- impacts 35 75 % of men who have diabetes.
- Men with diabetes are likely to develop the problem 10 15 years earlier than men without diabetes. Over the age of 70, there is a 95% likelihood of facing difficulties with erectile function.
- Women also experience sexual dysfunction at higher rates when they have diabetes. Up to 78% of women with diabetes will experience sexual dysfunction in the form of decreased lubrication, desire and arousal, dyspareunia, orgasmic dysfunction and sexual satisfaction. This rate increases if the woman has diabetes and depression and/or anxiety.

The following website has excellent up to date information about what ED is, the pathophysiology and the treatment options. https://urologywaikato.co.nz/patient-resources/erectile-dysfunction/

Often the person will not ask about ED issues. To help health professionals to become more comfortable at starting the conversation, prompts have been added to the common form.



There are a number of treatments for ED

- Glucose and BP management are the essential first steps of reversing erectile dysfunction.
- Counselling the basis of ED may be psychological, behavioural, or physical
- Exercise this does more than simply boost circulation, it can reduce weight, reduce depression and anxiety, it may increase testosterone, and it simply boosts how we feel about ourselves
- Oral medications there are many of these on the market. Some have a shorter half-life than others, they still require stimulation and arousal to achieve a positive result, and they have reduced in price considerably since coming off market. They are available without script from pharmacy, encourage the person to try them more than once before they say it doesn't work performance anxiety can reduce their effect the first time, and warn people about online herbal options particularly from overseas; the label claims are not always accurate
- Penile injections are often a useful option
- Vacuum devices
- Penile implant or prosthesis

MONITORING

- Refer to CHECKING rather than TESTING. Testing suggests a pass or fail. There are no "wrong" answers to the monitoring as it is investigative, not accusatory.
- People prescribed Metformin, Galvus, Jardiance or Trulicity are not at risk of hypoglycaemia unless
 also taking insulin or a Sulfonylurea and do not "need" to monitor their blood glucose from a
 safety aspect.
- However, self-monitoring of blood glucose (SMBG) is the only way the person with diabetes (PWD) and their health care professional know what is happening with their diabetes. The more checking being done the more information they (we) have.
- People can learn more about their own diabetes by looking at how their blood glucose respond to what they eat /do/feel/medication use.
- Short periods of intense checking 3-4 days are achievable by most and gives you the information that is required to make decisions on medication options.

Most people will be somewhere on the following continuum:



Pre breakfast, pre bed 1 day a week

Pre and 2-hour post meals (6) Plus pre bed (7) plus 2 am (8 checks)

- Encourage checking even when levels are not great. Identifying the problem times of day means you can work towards fixing the problem
- Consider getting people to write their results in a logbook/download results/use an app such as https://www.mysugr.com/en/ at home as they are looking for a pattern
- Discuss BGL patterns what can you see and what does this mean? CHO consumption, exercise and medication
- Use a food/exercise diary with blood glucose check results alongside for a 4-day snapshot (see pages 26/27 and photocopy this for your patient to use)

Paired checking and overnight hypoglycaemia

- If starting a new insulin or concerned about <u>overnight hypos</u> (people often sleep through these) ask the person to check at 2am.
 - Clues to overnight lows HbA1c is low, but daytime glucose levels are high, >0.5 units/kg basal prescribed, waking with nightmares or sweaty, elevated results in the morning (there can be a rebound like effect), or simply that something just does not make sense.
- Paired checking is when people check before and then 2 hours after a meal.
 The thinking is that if there is a postprandial rise of more than 3mmols, reassess CHO load of the meal. Is it excessive or reasonable? Or introduce/increase medications that support insulin secretion (Sulfonylurea, rapid acting insulin). Consider dietitian referral.

Whichever meter the person uses, the correct technique is important.

Wash hands prior to testing

Keep meter in upright position



Insert the square end of a CareSens N test strip: the meter will turn itself on.



Apply blood to the test strip.

Only 0.5 µL is required (that's a very small amount).



After 5 seconds the result is displayed.

Remove the test strip and the meter will turn itself off.

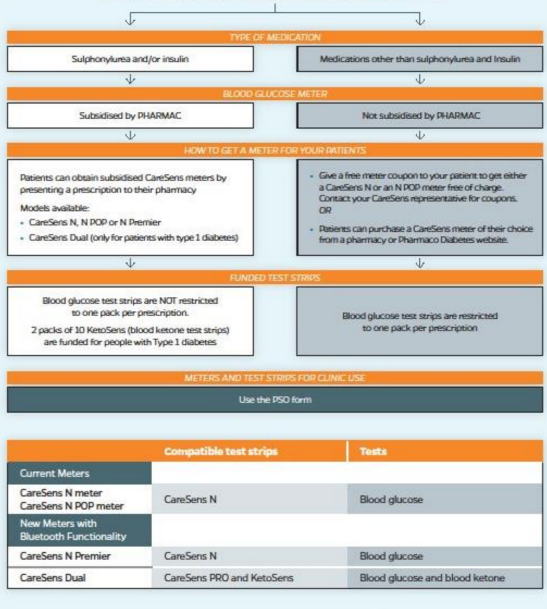
Easy.

Battery—CR2032 Lithium battery for the CareSens range of meters are free from your local pharmacy.



A guide to the availability of CareSens meters and test strips for newly diagnosed patients.

Patients newly diagnosed with diabetes



Pharmaco Diabetes Training Portal

The Pharmaco Diabetes Training Portal is a convenient online training platform for healthcare professionals. Simply go to the portal, register your interest and start learning at a time and place that works for you. Full of information and videos, the portal will help you become expert in using the CareSens blood glucose monitoring systems.

www.PharmacoDiabetes-training.co.nz

Lancets are not free but please ensure that people are changing them because when they are blunt, they often cause more pain and can lead to resistance to self-monitoring of blood glucose. One lancet is good for 12 finger pricks.

		and the second second	CareSens N	
	CareSens N	CareSens N POP	Premier	CareSens Dual
Features	555	55-	57	<u> </u>
Test	Blood Glucose	Blood Glucose	Blood Glucose	Blood Glucose and Blood Ketones
Compatible test strips	CareSens N	CareSens N	CareSens N	Care Sens PRO For blood plucose testing
	A		# N	Returbers 1
Bluetooth data transfer to SmartLog app	NO	NO	YES	YES
Manual data entry to SmartLog app	YES	YES	YES	YES
Data download to Windows and Mac computer	YES	YES	YES	YES
Illuminated numbers	NO	YES	YES	YES
Meal Flags	Pre and post meal	Pre and post meal	Fasting, pre and post meal	Fasting, pre and post meal
Memory	1000	1000	1000	1000
Averages	1, 7, 14, 30 and 90 days	1, 7, 14, 30 and 90 days	1, 7, 14, 30 and 90 days	1, 7, 14, 30 and 90 day
Strip ejector	NO	NO	YES	YES
Ideal for	People with diabetes looking for a simple easy to use meter to test blood glucose levels	People with diabetes looking for a compact and discrete meters to test blood glucose levels	People with diabetes needing advanced meter with Bluetooth functionality and bigger screen	People with diabetes for testing blood glucose and blood ketone typically people with type 1





www.PharmacoDiabetes.co.nz

Always read the label. Follow the manufacturer's instructions, and the advice provided by your healthcare professional.

Pharmaco (NZ) Ltd, Auddland. TAPS DA1827IM.

For more information on CareSens meters and test strips contact Pharmaco Diabetes on support@pharmacodiabetes co.nz or 0800 GLUCOSE (0800 45 82 67)

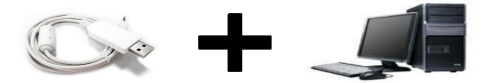
For more information on eligibility criteria please refer to the pharmaceutical schedule.

Pharmaco 0800 458267

Ensure you have enough replacement meters—use MPSO and the vouchers available from your Caresens rep. Mob. 0211959634 or 0800 glucose.

Smartlog – downloading meters

This is really useful for helping to adjust medication to BGL profiles



The software is free and the cord you can get free from the rep.

Patients can also buy their own + download the free software on the website

Olivia Solomons Territory Manager-Diabetes PH: 021 1959634 olivia.solomons@pharmaco.co.nz

https://www.pharmacodiabetes.co.nz/smartlog-data-management/smartlog-software/





There are some very helpful videos on the website to guide you or your patient set up functions on the meter https://pharmacodiabetes.co.nz/helpful-videos/

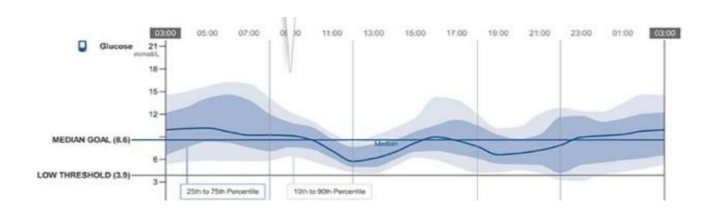
CareSens N Voice

- With its talking function, the N Voice makes checking easier for those who are visually impaired.
- In addition to the talking function, the N Voice has a larger screen and numbers.
- The N Voice comes with a test strip ejector for the easy removal of a used strip.





The **FreeStyle Libre2** Flash Glucose Monitoring System is a glucose monitoring device used to detect trends and track patterns without the need to finger prick. The system is not funded for type 2 diabetes, and the sensors are around \$118 (delivered) each. They last 2 weeks and can be blue toothed to a compatible phone. If the phone is not compatible, scanners can be purchased from the company. People can use them continuously or use them when they want more in-depth picture of their BGL, or when medication is being titrated. Ask your PHO diabetes lead if you are interested in trialling a patient on a Libre2.



More information can be found at Mediray New Zealand

https://www.mediray.co.nz/freestyle-libre/freestyle-libre-2-flash-glucose-monitoring-system/



Ketone Testing

A bit of Pathophysiology

When there is not enough insulin, the fat cells keep releasing fat into the circulation, and the liver keeps making more and more ketones and ketoacids. The rising ketoacid levels make the blood pH too low (acidotic), which requires immediate medical attention.

A person with Type 1 diabetes has the highest risk of developing ketoacidosis.

However, ketoacidosis also can occur in someone with Type 2 diabetes if there is a major increase in insulin resistance (<u>such as infection or treatment with steroids</u>) or reduction in insulin release from the pancreas.

People are most likely to develop ketones when unwell.

If a person with Type 2 Diabetes presents to general practice and is physically unwell, it is useful to do a ketone test. You can get Ketone tests on MPSO and they work with the CareSens duo meter, particularly if they are on SGLT2i (Jardiance).

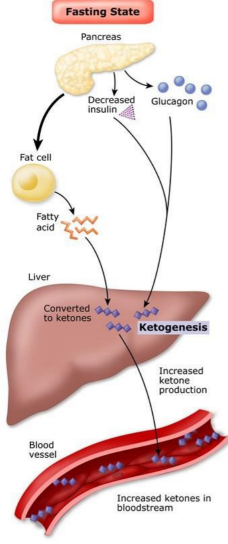
If unwell, someone taking Jardiance can have a glucose in the normal range but still have ketosis. The Jardiance prevents them passing ketones via the urine, so you must do a blood ketone test using the DUAL meter.

Ketone blood testing is the preferred method for assessing the presence of ketones during times of sickness, as they are more accurate and timelier than urine tests. When using a DUAL meter results will fall into one of the following three ranges:

- Below 0.6 mmol/L: Normal range
- 0.6 to 1.5 mmol/L: This may develop into a problem if not treated
- Above 1.5 mmol/L: Readings above 1.5 indicate a greater risk for developing Ketoacidosis (DKA).
 Discuss with GP and consider acute sick day management. Consider referral to hospital

Regularly check the expiry date of ketone strips as they have a short life

Ketone Production by Liver During Fasting Conditions (Ketosis)



NON-INSULIN MEDICATIONS

Common Oral Hypoglycaemic Agents

Full information can be found on the New Zealand Formulary (nzf.org.nz). Prescribers need to be aware that the following pages are for a *quick reference* only.

Metformin hydrochloride

Metformin is the first line of treatment and sensitises the patient to their own insulin. Initially 500mg 1-2 times daily, **increased gradually as tolerated** to 1.5-2g daily in divided doses; there is very little clinical benefit over 2g/day.

If you find people have had a GI upset from Metformin, encourage them to try again with a 250 mg dose once a day. Increase this very slowly. It doesn't matter how small the dose is and how slowly you increase it, there will still be some benefit!

Take it up as far as they can until they get GI upset again and then pull it back slightly and you have found their tolerated dose.

Patient Advice:

- Take with or just after food or at a meal to reduce side effects.
- Contact your doctor immediately if symptoms such as increasing drowsiness, loss of appetite, nausea, vomiting, abdominal pain, cramps, fatigue, diarrhoea, unexplained weight loss, muscle pain, and non-specific abdominal upset.
- Metformin may induce ovulation, so discuss contraception with women of childbearing age.

Renal Impairment:

- KDIGO now recommend that 2 g per day is safe when the eGFR is > 45 mL/min.
- eGFR 30-45 1000mg maximum daily dose
- eGFR 15-30 500mg maximum daily dose
- eGFR < 15 avoid.

Remember:

Withdraw or interrupt treatment in those at risk of tissue hypoxia or sudden deterioration in renal function, such as those with dehydration, severe infection, serious trauma, shock sepsis, acute heart failure, respiratory failure, or hepatic impairment, or those who have recently had a myocardial infarction - increased risk of Lactic acidosis.

STOP taking temporarily if transient tummy bug, prior to surgery or contrast dye procedure

(see page 77 - sick day mangement and 82 - planned surgery).



SGLT2inhibitors – Empagliflozin Jardiance

90% of glucose is reabsorbed from the urine back into the blood stream by Sodium Glucose cotransporter 2 in the proxmial renal tubual, along with sodium and water.

If this process is blocked or inhibited, then glucose and fluid loss in the urine is increased.

SGLT2inhibitors do just that, lowering blood glucose and blood volume.

Your patient will meet special authority criteria if they have type 2 diabetes with an HbA1c > 53 mmol/mol despite at least 3 months of regular use of metformin and/or an alternative glucose lowering therapy, is not on a funded GLP1RA (i.e. dulaglutide) AND any of the following:

- has diabetic renal disease (ACR > 3 mg/mmol and/or eGFR < 60 mL/min) OR
- has known cardiovascular disease (any ischaemic heart disease, cerebrovascular event, peripheral vascular disease, congestive heart failure or familial hypercholesterolaemia) OR
- has a 5-year cardiovascular disease risk > 15% OR
- has a high lifetime cardiovascular risk due to onset of diabetes in childhood or as a young adult OR
- is of Māori or Pacific ethnicity

Preferred 2nd line agent after metformin if renal disease and/or heart failure predominates

Doses:

Either once a day as Jardiance in a 10mg or 25mg tablet.

Or twice a day mixed with Metformin as Jardiamet

5mg / 500mg

Or

5mg / 1000mg





12.5mg / 500mg Or 12.5mg / 1000mg

Benefits of SGLT2i

- Reduction in HbA1c of 5 11 mmol/mol
- Hypoglycaemia is rare, but consider reducing insulin doses if HbA1c < 65
- Modest weight loss
- Mean reduction in systolic and diastolic BP of 1 6 mmHg
- Reduced progression of diabetic renal disease
- Reduced CVD events
- Reduced death from heart failure

Side effects of SGLT2i

- Lower blood pressure consider review of BP meds
- Urinary tract infections & genital thrush, which is usually mild and short lived, but patients need advice on genitourinary hygiene and to seek help if concerned.
- Consider back pocket script for thrust treatment if past Hx of thrush

If your patient is unwell with D&V they are at risk of DKA.
They should come to the practice for a ketone check. You must have a CareSens Dual meter and KetoSens test strips available.



GLP1RA – Dulaglutide (Trulicity) Liraglutide (Victoza)

Glucagon Like Peptide 1 Receptor Agonist is the main incretin produced in the small bowel. The mechanism of action of its receptor agonist is to act on the pancreatic islet cells to increase glucose dependent insulin secretion, decrease glucagon secretion and reduce hepatic glucose production. It also acts on other organs to slow gastric emptying, reduce appetite, increase cardiac output and increase loss of sodium.

Due to a worldwide shortage of supply, refer to the PHARMAC website for the latest on availability and SA criteria https://pharmac.govt.nz/medicine-funding-and-supply/medicine-notices/dulaglutide . Best practice does indicate that both can be prescribed, so if the patient chooses to self-fund one of the 2nd line therapies, the empagliflozin is considerably cheaper than the dulaglutide or liraglutide.

Preferred 2nd line agent after metformin if cerebrovascular disease predominates, especially if overweight or obese

Clinical benefits

- Mean ↓ in HbA1c of 6-15mmol/mol
- ↓ Weight
- Mean ↓ in systolic BP 2-3mmHg
- Mean ↓ LDL 0.08mmol/L
- ↓cardiovascular events (MI, CVA, CVD death)
- Likely \downarrow progression to diabetic renal disease

Adverse effects

- Gastrointestinal (nausea, vomiting, diarrhoea) usually mild and reduces with time
- Injection site reactions
- Antibody formation to GLP1RA
- Pancreatitis caution recommended if previous pancreatitis

Anticipate and prepare for adverse effects

Up to 30% of people suffer nausea when starting dulaglutide or liraglutide. Reassure your patient that the GI symptoms are expected, usually mild, <u>will reduce within 2 or 3 weeks</u> and are less if they:

- Have adequate hydration
- Eat smaller meals stop eating when you feel full
- Avoid fatty or spicy food
- Reduce smoking and alcohol intake
- Don't eat within 2 hours of bed

GLP1ra may induce ovulation. Discuss contraception

DULAGLUTIDE (Trulicity)

- Dose 1.5mg weekly injection (max dose
 4.5mg per week, or 3 injections
- Pre-loaded single use pen. Dispose of in sharps container after use.



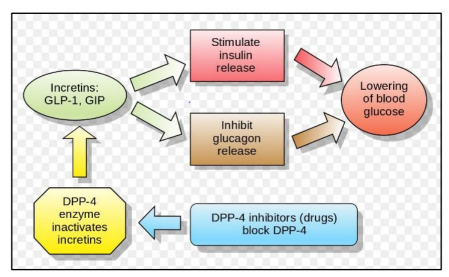


LIRAGLUTIDE (Victoza)

- Dose start at 0.6mg per day for a week, increasing step wise to 1.2mg per day for a week and then 1.8mg per day.
- Prescribe 4mm needles and change daily, provide sharps container.
- Prefilled pen lasts approximately a week.

DPP4inhibitors – Vildagliptin Galvus

<u>Dipeptidylpeptidase</u> is an enzyme that deactivates GLP-1 (glucagon-like-peptide-1) and GIP (glucose-dependent insulinotropic polypeptide). After eating a meal these hormones inhibit glucagon release and increase insulin production, they also delay gastric emptying, all of which lower blood glucose levels. DPP4 inhibitors block the enzyme DPP4 and enables the GLP-1 and GIP incretins to work properly so improving blood glucose control



Galvus is the next 2nd line agent after metformin for those <u>not</u> meeting funding criteria for SGLT2i or GLP1ra

Galvumet which combines Galvus and Metformin is available in 50/850mg tablets and 50/1000mg tablets. This decreases medication burden as patients take less tablets for the same effect. Anecdotally, people intolerant to metformin often tolerate it well in this combined tablet.

- Galvus and Galvumet cannot be used in conjunction with GLP1RA (Trulicity/Victoza) but can be used with SGLT2i (Jardiance)
- Vildagliptin can be used at a dosage of 50mg once or twice daily. It should be reduced to 50mg if eGFR is less than 50mL/minute/1.73m2. It does not cause hypos by itself but used in combination with sulphonylurea or insulin hypoglycaemia can occur.
- Vildagliptin has been shown to delay the progression to requiring insulin.
- Gliptins are a class of medications that are weight neutral. They can cause a reduction of HbA1c between 7-12mmol after 12 weeks of treatment.
- There seems to be few side effects but the most common of these are nasal stuffiness, headache and dizziness which can occur in 6-9% of patients.

<u>Contraindicated</u>: do not use if liver enzymes (ALT or AST) are two and half times the upper limit of normal pre-treatment. The use of DPP4 inhibitors have been associated with an increased risk of pancreatitis although there is still uncertainty and debate around the strength of this association but probably best to avoid in this group of patients.

A very comprehensive Bpac article can be found here: https://bpac.org.nz/2018/vildagliptin.aspx

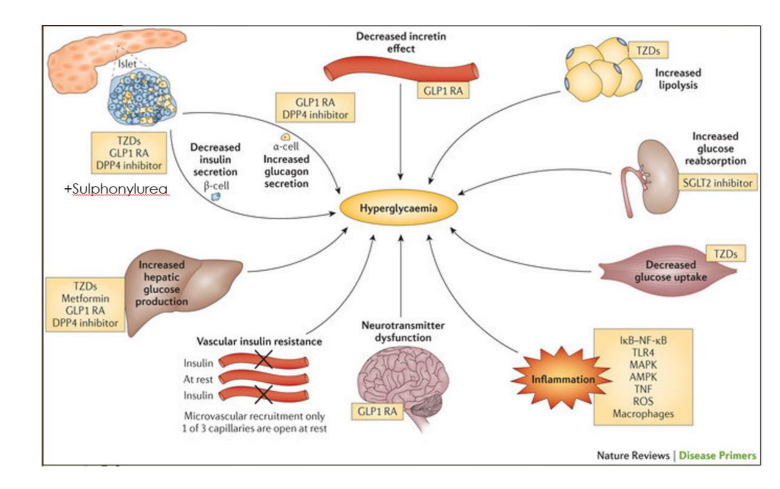
Other funded 3rd line oral agents

A snippet of less used/available oral medication options.

Find all the up to date information at: https://t2dm.nzssd.org.nz/Subject-21-Non-insulin-medications

Class	Name	Background Information		
		Stimulates the pancreas to produce more insulin		
		• Effectiveness diminishes over time as βcell function declines		
		Take with breakfast or first meal of the day. If BD dose, take second		
Sulphonylurea	Gliclazide	dose with evening meal		
	Glipizide	Can cause hypoglycaemia. Patients need education about hypo		
		management and a funded glucose meter		
		Can lead to weight gain		
		Do not use in conjunction with rapid acting or mixed insulin		
		Contraindicated: pregnancy, renal and hepatic impairment. Nation calls make a consisting to inculing.		
		Makes cells more sensitive to insulin		
		Shown to decrease cardiovascular outcomes		
		Beneficial with some mild MAFLD, high TG, or low HDL		
		Caution in patients with renal dysfunction/liver failure		
Thia-	Pioglitazone	AVOID in patients with heart failure		
zolidinediones		• 2.3-4.9kg weight gain		
(TZD)		 Side effects: Peripheral oedema, expansion of plasma volume, increased risk of anaemia, heart failure 		
		• <u>Contraindicated</u> : History of heart failure, bladder cancer, osteoporosis – fracture risk in women if used long term		
		 Only continue beyond 6 months if there has been a reduction of 5mmol/mol or more in HbA1c 		
		Taken orally and reduces the amount of glucose absorbed in the small intestine by blocking the a-glucosidase enzymes.		
		Should be taken with a meal.		
		Can be used as first line or added to any of oral anti-diabetic meds		
a-Glucosidase	Acarbose	 Hypoglycaemia can occur if used with insulin/sulfonylurea - patients should consume glucose not sucrose if hypos occur 		
Inhibitors		Flatulence, soft stools and diarrhoea are common side effects		
		Abdominal distension, pain and hepatitis have been reported		
		 <u>Contraindicated</u>: in pregnancy, people with hepatic or renal impairment, have inflammatory bowel disease, history of intestinal obstruction/hernia, previous abdominal surgery or GI disorders with malabsorption 		

A visual overview of non-insulin medications mechanisms of action



Each of these different medications has a different mode of action.

Not all medications are available or fully funded in New Zealand. People emigrating into New Zealand may be using medications which they do not meet the funding criteria for here, and they may have to pay. They may be on a similar medication, and if they do meet the criteria, they might like to change to the funded version e.g. Dapagliflozin is not funded in New Zealand, but Empagliflozin is.

If a person is already on insulin, you can still initiate oral medications to achieve the agreed HbA1c target. Your patient may have started insulin prior to these newly funded medications being available. Introducing new non-insulin medications may be an opportunity to decrease insulin doses, reducing hypo risk.

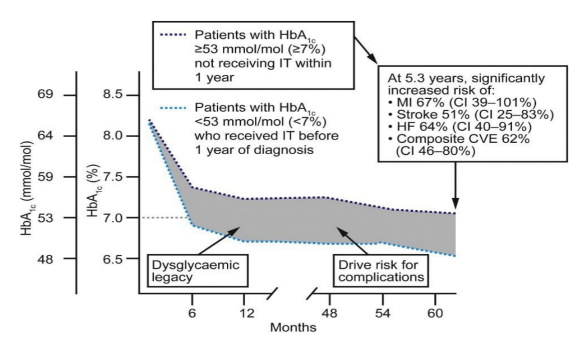
INITIATING INSULIN

When starting a person on insulin have a clear plan of what you are aiming to achieve; the agreed HbA1c target, know that they are already on maximum oral medication, and that the person has had some time to talk about their concerns. Have a check list, clear documentation, and a follow up strategy to ensure the correct dose is obtained. The process takes time and the whole practice needs to support the additional time and effort this takes.

The following chapter starts with two check lists. One is the process involved, the next is an education check list for the patient. Then each of the points on the education check list is explained in detail. This may be completed in one session, but in reality, only a small portion of what is taught is retained. You will need to create multiple opportunities to reinforce the information in the short term, but also over the years. The DAR is always a great opportunity to ask questions related to insulin administration such as hypo management or sharps disposal.

The earlier the insulin is started, the more of the remaining beta cell function is preserved.

This is a sobering graph showing the consequences of delaying insulin treatment (IT) by 1 year taken from Primary care diabetes review regarding clinical inertia.



If you need any support or further education regarding insulin initiation,

contact Lauren Southgate on 027 333 8895 your Pinnacle diabetes lead.

Have a plan:

Before—During -After

<u>Before starting insulin – consider these factors:</u>

- What other things could be causing hyperglycaemia? Consider food, fluids, exercise, stress, medication, lack of sleep, unwell, illness.
- Dietitian review to prepare for mixed or bolus insulin
- Check understanding of why control is necessary.
- Do they take their oral meds?
- Are the orals at maximum dose for that patient?
- What are their blood glucose levels like? logbook profile. No checking? No insulin!
- Address their concerns about starting insulin.
- Script: Insulin (plus 2 units for priming)
 - Needles BD microfine 4 mm / 5 mm
 - Test strips if required.

Your practice needs to have a clear pathway for getting the initial dose of insulin to correct level.

Either prescriber led titration, or patient led titration can work.

During insulin initiation - consider these factors:

- You will need a 45 60 minute appointment.
- Use check list.
- Consider having check list for patient as well.
- Company handouts.
- Principles of adult education (don't 'tell' patient what to do!)
- Lots of opportunity for hands on with the device.
- Consider having first dose while in clinic.
- Key information vs information overload. Dose/pen/hypos.
- Clearly arrange next contact moment.
- Document if you haven't recorded then it didn't happen.
- Claim Insulin initiation through primary options.

After the initial education session – consider these factors.

- Phone call, email, face to face.
- Blood glucose levels.
- One 'topic' e.g., hypos "so tell me what it would feel like, what have you got prepared to treat...."
- Document.
- Insulin adjusted with GP.
- Next contact arranged.
- Exit appointment ensure all information covered and documented.
- Next blood test arranged and encouraged to make contact if anything changes.

Checklist

Using a check list helps you and the patient know what to expect and provides a framework for the education process. The following is one example.

Starting Insulin

Patient name:	Date:	
---------------	-------	--

Checking

When to check, how to check, how to record.

Increase if unwell, check if hypo, looking for patterns.

Insulin regimen

Which insulin, what dose, and when to administer.

How to use injection device and importance of priming.

How to administer the insulin.

System in place for titration.

Sharps disposal.

Storage of insulin.

Hypoglycaemia advice

How to recognise, manage and treat hypos.

Formulate individualised plan of action – what and where will their hypo kit be.

Discuss hypo prevention.

Oral medications

Any changes?

Lifestyle advice

Maintaining healthy weight by healthy eating and exercise and reinforce advice from the dietitian.

Risks with alcohol consumption and insulin.

Driving

Ensure understanding of their responsibility to maintain a reasonable level of glycaemic control when driving. Particularly how to minimise their risk of hypos.

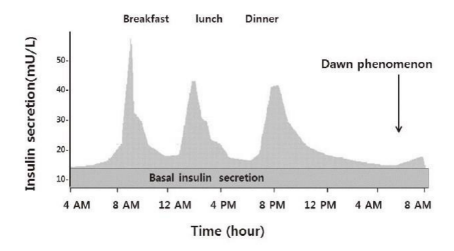
If vocational driver on insulin – Ensure control is adequate for safe driving. If required, or you are uncertain, refer for specialist advice.

Next contact time:	

Practice Tip: DOCUMENTATION:

Set up a key word that mimics your education check list. It is then simple to document what you have covered today and what needs to be covered next time.

Background information



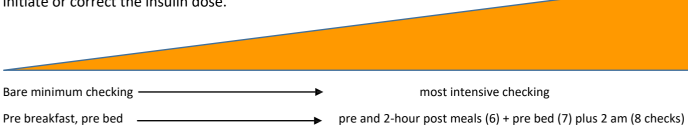
This is the insulin profile when we do not have diabetes and we eat 3 meals a day. When people have diabetes, and their blood glucose control deteriorates the aim of treatment is to mimic this natural profile, without causing hypos.

Initially the natural function of the pancreas is supported by adding in a basal insulin. The aim being to reduce the overall BGL, assuming that the beta cells can still make enough post prandial (after food) insulin on their own. Over time, the ability to make post prandial insulin diminishes and we need to also add a rapid acting insulin that matches the food intake (called a bolus dose). This can be as a separate dose of rapid acting insulin, or a premixed insulin.

Blood glucose checking when starting/changing insulin

Refer chapter 5 on *monitoring* for more information.

While insulin is being initiated and titrated to the correct dose, a commitment to some glucose checking is required. Without knowing what the blood glucose pattern is for the individual, we cannot initiate or correct the insulin dose.



2 am checking

If starting a new insulin or concerned about overnight hypos (often people sleep through these) ask the person to check at 2 am. Clues to overnight lows – HbA1c is lowish, but daytime checking high. Waking with nightmares or sweaty, elevated results in the morning (there can be a rebound like effect), basal dose >0.5 units/kg, or simply that something just does not make sense.

Paired checking

Refers to checks before a meal and 2 hours after a meal. The thinking is that if the rise between the two checks is more than 3mmols, a rapid insulin is required with the meal (see page 41).

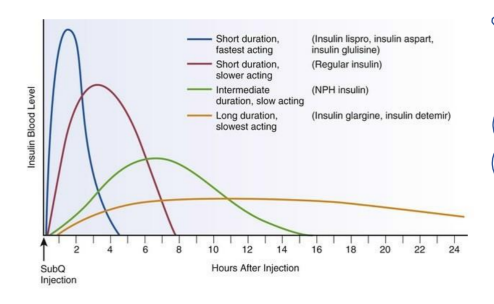
Insulins to choose from

Three companies provide insulin – Sanofi, Novo Nordisk and Lilly. Get to know the reps and keep up to date with their handouts. There are a limited number of insulin options – Long acting, intermediate acting, rapid acting and several premixed options. There is also short acting – but rarely used in primary care. Most clinics are very familiar with 3 – 4 insulins and get to know how they work and when one is more appropriate than the other. Be aware there are more options than this and be open to trying different types. One insulin option will not work for every person. The trick is to look at the blood glucose profile and fit the insulin that best matches, while also considering the person's lifestyle and the goals of management. Consider that a person may need one insulin in the morning and a different insulin regimen in the evening; or weekday at work is one regimen and weekends they take something different. See Appendix 3 (page 89) for a guide to help choose appropriate insulins for your patient.

Insulin regimens

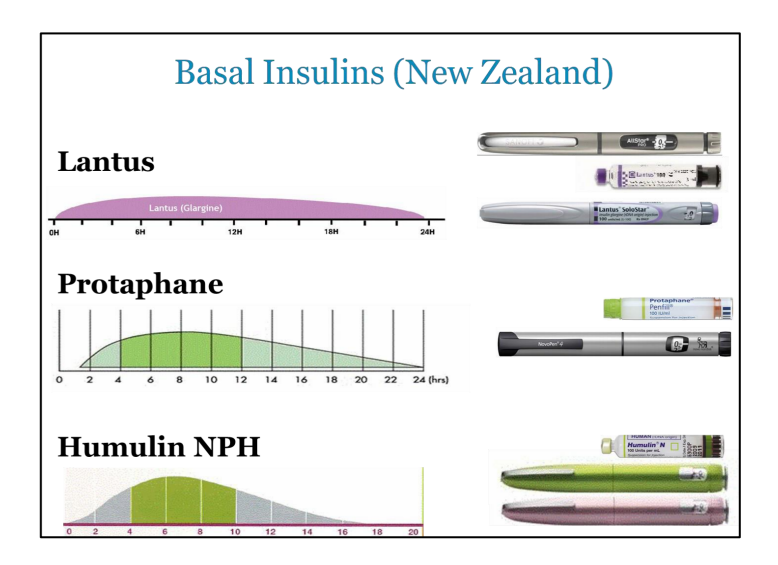
Simplest most complex

Long acting daily	Intermediate daily and BD	Premixed	Basal bolus
Once day, doesn't intrude into lifestyle (often taken at night) Dose is same time every day and does not need to be taken with food.	Works well for people with high BGL at one end of day but not the other. Simple. Easy to progress from once a day to twice.	Gives some post- prandial control. Once twice or three times a day.	Most flexible, can achieve great results pre and post prandial Consider one, two or three bolus doses depending on need
Gives no post prandial control (relies on some residual insulin supply) Impact on BGL may not cover 24 hours	Gives no postprandial control.	Best for people whose lifestyle is routine. Less flexibility, but also less work. Taken just before or with meals.	Works best if checking pre (and post) meals, CHO counting and awareness of activity impact.



Take time to discuss with the person the difference in insulin regimens.

You may be surprised what they choose.



Rapid Insulins (New Zealand)

Peak at 1-3 hours, lasts up to 5 hours

Insulin type	Brand name	Manufacturer	Nature	Appearance
Insulin lispro	Humalog®	Lilly	Analogue	Clear
Insulin aspart	NovoRapid®	Novo Nordisk	Analogue	Clear
Insulin glulisine	Apidra®	Sanofi	Analogue	Clear



Premixed Insulins with analogue (rapid)

Includes rapid-acting insulin.

Insulin type	Brand name	Manufacturer	Nature	Appearance
Lispro 25%/lispro protamine 75%	Humalog® Mix25™	Lilly	Analogue	Cloudy
Lispro 50%/lispro protamine 50%	Humalog® Mix50™	Lilly	Analogue	Cloudy
30% rapid-acting / 70% protaminated insulin aspart (rys)	NovoMix® 30	Novo Nordisk	Analogue	Cloudy



The following insulins are not used as much but are still available in New Zealand. They contain short acting, or human insulin as opposed to analogue insulin (rapid acting) Note—human insulin is manufactured in a laboratory using recombinant DNA technology— not from humans per se!

Short acting insulin with human component



Peak at 2-5 hours, lasts 6-8 hours

Insulin type	Brand name	Manufacturer	Nature	Appearance
Neutral	Actrapid®	Novo Nordisk	Human	Clear
Neutral	Humulin® R	Lilly	Human	Clear

Any insulin that is premixed (Novomix30 or Humalog Mix), or cloudy in nature (Humulin NPH or Protaphane), needs to be gently resuspended prior to injection. Use a <u>rolling</u> or similar action not a shake, which will introduce bubbles.

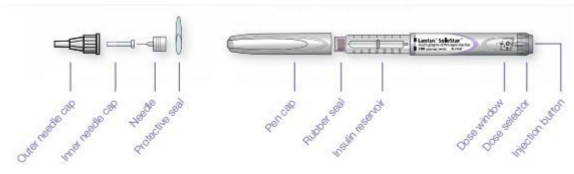
Devices

Consider different devices – <u>disposable</u> (best if there is less dexterity, convenient), or <u>non-disposable</u> (environment?), what does the patient prefer. Familiarise yourself with the pens prior to starting a patient on insulin. A well-resourced kit such as the one below means you will have all the gear needed in one place in your clinic.

Practice tip:

A well-resourced demonstration kit such as this ensures that you have all the demonstration pens, needles, cartridges and the dummy tummy in one place.





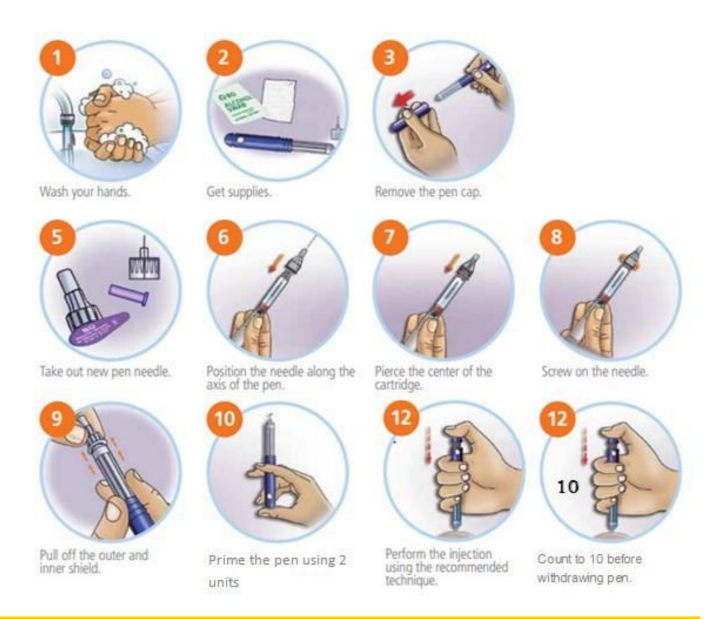
Follow manufacturer's instructions. https://www.lantus.com/-/media/ems/conditions/diabetes/brands/lantus/consumer/lantus-solostar-pen-guide.pdf

Make sure you know which insulin goes with which pen.



Technique

When teaching a patient to use insulin, it is a good idea to get the "injection part" done early in the session, otherwise they are so anxious about the injection that they hear very little of the education you provide. Once they see it's not as bad as anticipated, they relax and are in a better space to hear and learn.



"Priming the pen" means dialling up 2 units and squirting into the air to make sure there are no air bubbles in the pen or needle. The patient then dials up the dose and injects into their thigh or abdomen. The prescription needs to allow for priming or they will run out of supply before the end of the month.

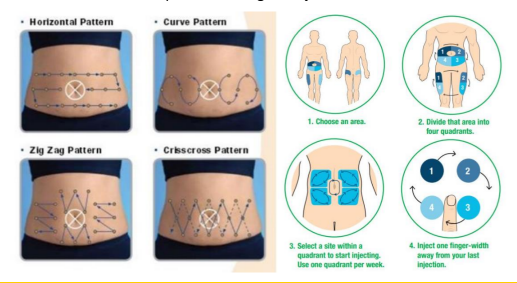
Any insulin that is premixed such as Humalog Mix50, or cloudy in nature such as or Protaphane, needs to be gently resuspended prior to injection. Use a <u>rolling</u> or similar action not a shake, which will introduce bubbles.

Starting Basal Insulin

Insulin can be used in conjunction with oral medications, although those that are insulin secretagogues (e.g. sulfonylurea) may need to be down titrated or switched to bolus insulin if hypoglycaemia occurs.

- Start isophane (Protaphane or Humulin NPH) or glargine (Lantus) insulin at
- o 0.1 units/kg daily if HbA1c < 64 mmol/mol or BMI < 18 or elderly or renal/liver failure
- o 0.2 units/kg daily if HbA1c > 64 mmol/mol and BMI >18
- Start with a nocte dose to counteract hepatic gluconeogenesis and target fasting blood glucose.
- Monitor fasting blood glucose levels, and educate on how to manage hypoglycaemia (page 71)
- If 3 consecutive fasting blood glucose levels are > 7mmol/L then increase the dose by 10% or 2 units every 3 days (see page 67 for more on titration)
- Stop up-titration of basal insulin if;
- O Hypoglycaemia occurs (< 4mmol/L), OR
- Fasting blood glucose is < 7mmol/L, OR
- Dose reaches 0.5 units per kilogram per day
- Basal insulin can be injected into the thigh or abdomen
- Prescribe 4 or 5mm needles for optimal absorption, which should be changed daily
- Rotating the site of injection is important to avoid build-up of scar tissue (lipohypertrophy) which results in poor absorption over time

Examples of rotating the injection sites:



Treatment intensification should be considered for patients who have not reached their HbA1c target after 3 months of using basal insulin, despite achieving fasting levels of <7mmol/L</pre> and/or they are taking a dose of 0.5units per kilogram per day.

They may need bolus or mixed insulin...

Starting Mixed (biphasic) Insulin

Mixed insulins are the best option for some one who requires insulin to improve glycaemic control after meals, and:

- Does not need flexibility for work patterns or exercise
- Eats similar meals at similar times each day
- Is not likely to need rapid intensification of insulin therapy
- Has reduced dexterity, cognitive ability or requires suppervison/support with medications
- Prefers not to regularly monitor blood glucose levels
- Prefers fewer injections daily

When converting from basal only insulin to mixed insulin if the person generally eats one big meal per day;

- Convert the total daily dose of basal to premixed insulin and inject before the largest meal
- Monitor BGLs before and 2 hours after meals (paired checking, page 41)
- Increase dose by 10% if
- 3 paired checks indicate a rise of > 3mmol/L after meals AND
- o 3 fasting levels are > 10mmol/L

When converting from basal only to mixed insulin if the person usually eats more than one meal a day;

- Convert the total daily dose of basal to premixed insulin and inject half before breakfast and half before the evening meal
- Consider a different ratio (e.g. 1/3 with breakfast, 2/3 with dinner) if one meal is larger than the other
- Monitor BGLs before and 2 hours after breakfast and dinner (paired checking, page 41)
- o Increase breakfast dose by 10% if 3 paired checks indicate post prandial rise of > 3mmol/L AND 3 predinner BGLs are > 10mmol/L
- Increase dinner dose by 10% if 3 paired checks with dinner indicate a post prandial rise of > 3mmol/L AND
 3 pre-breakfast BGLs are > 10mmol/L

Rotate the injection sites every time to avoid lipohypertrophy (lumpy bumpy under the skin)

Patients starting mixed insulin meet the referral criteria to the extended care team to see a dietitian to learn about eating a consistent amount of carbohydrate at meals. This will improve post prandial BGLs, lower HbA1c and help prevent weight gain.

Starting Bolus (Rapid) Insulin

Bolus insulin with meals is the best option for someone who requires insulin to improve glycaemic control after meals, and;

- Requires flexibility for work patterns or exercise
- Prefers a varied diet in terms of quantity and timing
- Will likely require rapid intensification of insulin therapy
- Has good ability to inject
- Is comofortable with monitoring blood glucose levels more frequently
- Is comfortable with more frequent injections 4 or 5 in total per day (basal + bolus)

"Basal Plus" is the addition of bolus insulin to the largest meal, while continuing the basal regimen.

- Start bolus insulin at 4 units or 10% of the basal dose
- Monitor BGLs before and 2 hours after the meal (Paired checking, page 41)
- Increase dose by 2 units every 3 days if 3 paired checks indicate a post prandial rise of > 3mmol/L
- Consider the early addition of bolus insulin at other meals if the paired checking indicates a post prandial rise of > 3mmol/L at the other meals
- Add correction doses to treat preprandial hyperglycaemia (see page 66)

Doses of basal insulin may need to be decreased to prevent hypoglycaemia, especially if HbA1c < 64mmol/mol

If the bolus dose is "right", a person will only need a correction dose occasionally. Correction doses are given when a persons blood glucose levels are unexpededly high before a meal. If someone is taking a correction dose at every meal or even every day then the bolus dose needs review and up titration from the prescriber (see page 67). The "right dose" is achieved when the blood glucose check 2 hours after the meal is consistently only 1 or 2 mmol/L higher than the pre meal level.

Patients starting bolus insulin meet the referral criteria for the extended care team to see a dietitian to learn about managing carbohydrate intake either through carbohydrate portions or carbohydrate counting. The dietitian will assess the most appropriate approach depending on the patient's health and maths literacy. This will improve post prandial levels, reduce the need for correction doses, lower HbA1c and help prevent weight gain.

Correction Doses

Rapid acting insulin can be used to "correct" high blood glucose levels when added to the bolus, or mealtime dose, of insulin. The only insulins that can be used for this are Novo Rapid, Humalog or Apidra. It is **NOT** suitable for basal or mixed insulins.

We ask our patients to check their blood glucose 2 hours after a meal as a safety measure, or to establish a pattern so we can determine if the bolus dose requires titration (page 69). It is **NOT** for the purpose of correcting a higher than expected reading. If someone requires correction most days or always at a certain meal, they likely require titration at the *previous* meal.

Rapid insulin is active for up to 5 hours after the injection. If a correction dose is given 2 or 3 hours after the meal, the bolus dose has not had a chance to work fully, and the additional dose is "stacked on top" of the initial dose. This can cause hypoglycaemia and is why it is important that bolus insulin is not given more than 3 times a day - at breakfast, lunch and dinner.

The correction calculation is based on the total daily dose of insulin

Total daily dose of basal + bolus insulin	Correction dose calculation
≤ 25 units/day	1 unit for every 4 mmol > 8 mmol/L
26 – 40 units/day	1 unit for every 3 mmol > 8 mmol/L
41 – 75 units/day	1 unit for every 2 mmol > 8 mmol/L
≥ 76 units/day	1 unit for every 1 mmol > 8 mmol/L

• An example of how to use correction insulin

- o If a patient is on 40 units of basal insulin nocte and 10 units of bolus insulin with meals their TDD is 70 units per day, so their starting correction is 1 unit for every 2 mmol > 8 mmol/L
- A table can then be provided for the combined doses of bolus + correction insulin at each meal e.g.:

Blood glucose level	Correction dose of insulin	Total insulin dose with meal
< 10 mmol/L	0	10 units
10 – 11.9 mmol/L	1	11 units
12 – 13.9 mmol/L	2	12 units
14 – 15.9 mmol/L	3	13 units
16 – 17.9 mmol/L	4	14 units
18 – 19.9 mmol/L	5	15 units
≥ 20 mmol/L	6	16 units

Rapid acting insulin is used to correct **PREMEAL** hyperglycaemia. It should not be injected between meals.

A patient using bolus insulin should only ever give 3 bolus injections per day.

Titration

- Ensure you have up-to-date contact details cell phone, landline, email, NOK.
- Make a time and date you will contact them for BGLs and to check how they are managing.
- Every phone call is an opportunity to reiterate the education you have given them. Pick one topic each time and go over it. Ask them for what they remember so you can ensure they did understand all the detail.

Typically, a person will be started on a small amount of insulin then the dose is titrated up to what they actually require. Talk with the person about what they may expect. This process may take some time and ongoing commitment to blood glucose monitoring.

Other than the doctor, only a nurse <u>prescriber</u> or pharmacist <u>prescriber</u> can change a prescription

- If the prscriber writes a maximum dose in the prescription, the nurse can support the patient to increase a dose up to that point.
- Patient can self-titrate at prescriber direction (using titration rates provided by company). Ensure that patient knows an upper limit to titration otherwise they may just keep going!
- A weight gain of 2 3 kgs is reasonably normal (not everyone gains weight especially if insulin is started earlier).
- Contact every 3 7 days until BGL within target. Don't take too long to reach ideal dose (risk is patient disengages). If the patient is quite insulin resistant, they may need large doses of insulin, however there is very little clinical benefit in greater than 0.5units per kilo body weight of basal insulin.
- Consider a percentage increase rather than a unit increase if the person appears to be needing quite a number of increases.

Always be aware that insulin doses may need to decrease if the person is experiencing lower than target blood glucose levels. Talk to the prescriber about this.

- Basal insulin needs to be given at the same time each day.
- Premixed and rapid insulin must be taken immediately prior to a meal.

If the patient is gaining weight or is regularly hypoglycaemic; they are on too much insulin.

Type 2 diabetes is not an acute condition – aim to commence and titrate insulin safely, but efficiently.

Be clear about the agreed HbA1c target with the patient. Tight control is not the aim for everyone.

Troubleshooting – I Help

Try this mnemonic if there is unexplained hyperglycaemia:

I HELP

I for Insulin

- ✓ is the correct insulin being used?
- ✓ Is the correct insulin in the correct pen?
- ✓ Is the insulin taken at the right time and sites rotated?
- ✓ Is the patient priming the needle and changing it regularly?
- ✓ Are doses of insulin being missed?
- ✓ Is mixed/cloudy insulin being mixed prior to use?
- ✓ Is the insulin being stored correctly?
- ✓ Is the insulin potent/expired/gone off due to extreme temperature change?
- ✓ Is the insulin being absorbed? Check the site for lumps and bumps.
- ✓ Is the correct needle length being used/ (4 or 5mm)

H is for Health status

- ✓ Is there underlying illness?
- ✓ Has a new medication been prescribed e.g prednisone, that changes glucose metabolism?
- ✓ Is there a change in renal function?
- ✓ Has there been a change in eyesight making dialling correct doses difficult?
- ✓ Is the patient dehydrated?
- ✓ Has the patient's weight increased lately?
- ✓ Check for ketones.

E is for Equipment

- ✓ Check glucose meter and insulin pen are working.
- ✓ Check strips/insulin expiry dates.
- ✓ Check storage of strips (not out of the container).
- ✓ Observe technique for checking BGL and administration of insulin.
- ✓ If on an insulin pump, contact the pump team.
- ✓ Chek patient is washing hands prior to checking BGL.

L is for Lifestyle

- ✓ Has there been change in diet e.g. a new food/drink or dietary strategy?
- ✓ Has there been a change in exercise habits?
- ✓ Check alcohol and drug consumption.

P is for Psychological issues

- ✓ Are there stressors that impact on the patient's ability to manage their diabetes?
- ✓ Are the prescribed doses of insulin and non-insulin medications being taken?
- ✓ Is there a fear of hypoglycaemia that leads to a reduced dose of insulin being administered?
- ✓ Does the patient have a needle phobia?
- ✓ Are there underlying mental health issues?
- ✓ Are doses of insulin being missed to control weight?

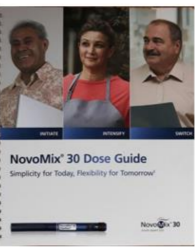
Intensification

There are times when a person needs a different insulin regimen.

- They started on a very simple regimen of one daily dose of insulin, but they no longer achieve adequate control on this.
- They require fewer doses of insulin (i.e. as people age, a basal bolus regimen that requires CHO counting and dose adjustment may be too complex for them to manage, or they simply don't wish to any longer).
- They are planning pregnancy and require excellent control of their diabetes in the preconception time.

Ensure that your clinic has up to date resources from each of the companies. Contact the representative if you have any questions.







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Current as of July 2024

Sharps containers

available in pharmacies for patients

Sharps containers are **ONLY** for sharps!

- Disposable pens can be thrown away—only the needle goes into the sharps container.
- Insulin cartridges can be thrown away—only the needle goes into the sharps containers.
- Plastic caps, band-aids, tissues ...do not belong in sharps containers
- Lancets can be disposed of in sharps container
- The aim is one small sharps container should last about a year!



Pharmacy

For those prescribed insulin or other injectable medications ¹ patients should use a plastic container no larger than a pump bottle and take that container to the pharmacy for disposal.



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¹ note: not for needle exchange programme sharps

Hypoglycaemia

- What does a hypo feel like? People need to be clear between the symptoms of hypoglycaemia and simply feeling tired/rundown. If someone feels the "symptoms" of a hypo, it is always better to check blood glucose levels before treating.
- Give written information.
- No driving for ONE hour after a hypo has been successfully treated.
- No driving for 24 hours after a severe hypo that has been treated by someone else because the person could not treat their own hypo.

Practice point:

Ask the person—"What have you got with you today if you needed to treat a hypo?" or "What have you got in the car?"

Appropriate treatment of hypoglycaemia will likely prevent severe rebound hyperglycaemia

Treatment of hypoglycaemia (blood glucose level < 4 mmol/L) is weight-based. If the **patient weighs > 70 kg they should eat 30 grams of rapid** acting carbohydrate and if the **patient weighs < 70 kg they should eat 15 grams of rapid acting carbohydrate**

- How to get 30 grams of rapid acting carbohydrate (halve for 15 grams)
 - i. 10 Dextro or vita glucose tablets or 6 BD glucose tablets
 - ii. 30 grams of glucose powder
 - iii. 6 teaspoons of sugar dissolved in water
 - iv. 350 mL of fruit juice or non-diet soft drink
 - v. 18 jellybeans
 - vi. 2 tablespoons of honey
 - vii. 3 tablespoons of jam
- viii. 2 Hypofit gels
- Wait 15 minutes and re-check blood glucose level
 - i. If glucose level < 4 mmol/L repeat steps above until glucose > 4 mmol/L

Waiting gives the glucose time to be absorbed

- Once glucose is > 4 mmol/L have a carbohydrate snack e.g. a slice of toast, 2 biscuits or crackers with cheese and recheck glucose levels in 30 minutes to ensure safe
- If the patient becomes unresponsive and is unable to take carbohydrate replacement an ambulance should be phoned for. Glucagon 1 mg may be administered intravenously or intramuscularly, but has reduced efficacy in type 2 diabetes and should not delay treatment with IV or buccal glucose.

Consider a medical alert bracelet https://www.medicalert.co.nz/net/products/

Hypo unawareness—this happens when someone has had so many hypos that their body no longer thinks it is unusual and forgets to warn them when their blood glucose levels drop below 4 mmol/L. This means they can have a blood glucose level of 2 or less in some cases, and yet feel absolutely normal. This puts them at extreme risk of going unconscious without warning. They can partially reverse this condition by keeping their blood glucose levels a little higher with NO hypos for a number of weeks or months. This helps 'reset' their body warning system so that they start experiencing symptoms of hypos when their blood glucose levels fall below 4mmols/L.

Insulin storage

- Insulin is stored in the fridge.
- Once opened store at room temperature (so the injection does not hurt as much) Dispose
 of opened insulin 28 days after opening.
- Check the expiry date before using.
- Look at the insulin—does it look like it normally does? Any particles, discolouration, crystal formation?
- Frozen insulin cannot ever be used—throw away.
- Do not keep insulin in the glovebox of the car—the temperature can rise too high and cook the insulin.
- Cooling wallets are designed for people travelling to hot climates and are activated by water
 no refrigeration needed. Each soak in water keeps the bag cool for 2 days.

The Frio wallet is available in different sizes and available here:

https://www.mediray.co.nz/diabetes/product-category/insulin-cooling-wallets/







Alcohol

Alcohol is very energy dense and it is easy to gain weight if drinking regularly.

Alcohol inhibits hepatic gluconeogenesis and induces hypogylcaemia whenever gluconeogenesis is required to maintain normal glucose levels. People with diabetes on Sulfonylurea or insulin should be warned about the blood glucose-lowering action of alcohol.

The recommendations for people with diabetes remain the same as for all New Zealanders.



Some tips from the American diabetes association:

- If you have diabetes, do not drink on an empty stomach or when your blood glucose is low, since your risk of low blood glucose increases after drinking.
- Don't skip a meal if you are going to drink. (If you use carbohydrate counting to plan meals, it is important to understand how the drinks you choose affect your blood glucose and often your insulin dose will need to be decreased if having more than one drink)
- Wear an I.D. that notes you have diabetes. If you are in a setting where people are drinking alcohol, hypogylcaemia may be mistaken for being drunk.
- For mixed drinks, choose calorie-free drink mixers like diet soda, club soda, diet tonic water or water.
- As with anyone with or without diabetes, do not drive or plan to drive for several hours after you drink.

Alcohol can cause hypoglycaemia shortly after drinking and for up to 24 hours after drinking. If you are going to drink alcohol, check your blood glucose:

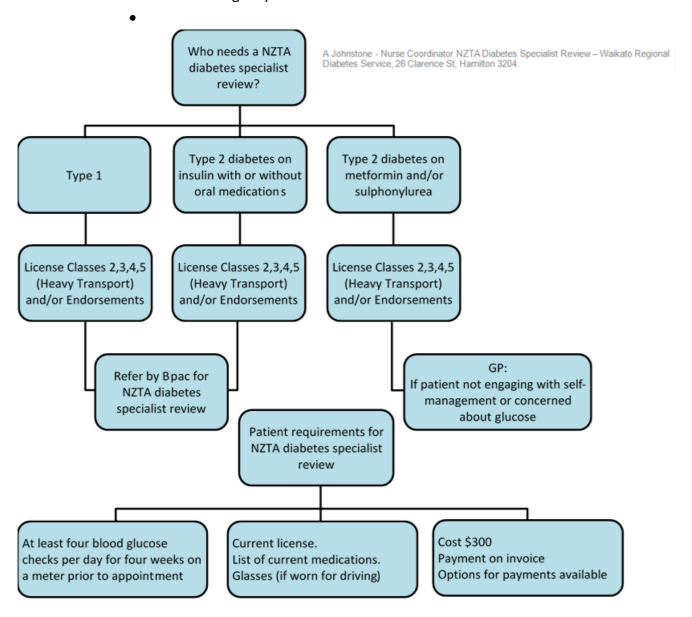
- Before you drink
- Before bed and throughout the night—needs to be over 8mmols before you go to sleep

If your blood glucose is low, eat something to raise it and be sure to check again before you go to sleep, and again over night to be sure it's not dropping too low.

Wondering if adding a glass of wine or beer might help lower your blood glucose if it is high? The effects of alcohol can be unpredictable, and it is not recommended as a treatment for high blood glucose. The risks likely outweigh any benefit that may be seen in blood glucose alone.

Driving

- The biggest risk is that a person has a hypo while driving causing an accident.
- Think—over 5 to drive
- Tips: Test prior to driving and regularly while on a long drive—every 2 hours is recommended.
- Keep your blood glucose meter with you while driving for evidence of regular testing
- Have a hypo kit in the car—nonperishable sweet items—glucose tablets and a longer acting CHO—muesli bar, crackers or similar.
- The NZTA advise that you should not drive for:
 - One hour after a mild hypo
 - 24 hours for a sever hypo (requiring some else's help)
 - 1 month if severe hypo while driving. An urgent review is required.
 - Your insurance may not be valid if you have an accident and have no evidence of monitoring on your meter



SICK DAY MANAGEMENT

This refers to periods of minor inter-current illness that requires changes to the persons usual diabetes self-management. It is important that patients are educated on how to deal with sick days and have an individualised plan and this should be revised at each annual review.



"Do you know your sick day rules?"

Key Concepts of sick day management

- Have a sick day management plan
- Never stop your usual insulin, doses may need adjusting
- Hourly fluids to reduce the risk of dehydration (1 glass/cup)
- Increase the frequency of blood glucose monitoring and blood ketones as necessary (1-hourly if ketones present, if not then 2-hourly if BG >15mmmol)
- Diabetes medication dose adjustment as necessary (type 2) and insulin adjustment (type 1 & 2)
- Metformin and Jardiance should be stopped for the duration of the illness
- Supplementary insulin as necessary (use individualised plan)
- Health professional number and name for assistance when sick—especially after hours.

HHS: high glucose levels can cause <u>hyperosmolar hyperglycaemic state</u> (HHS).

This condition is characterised by hyperglycaemia, hyperosmolarity and an absence of significant ketosis. The blood glucose is usually >30mmol with a raised osmolarity >320mOsmol, dehydration with no significant ketones/or acidosis.

There is insulin deficiency, and this condition can lead to coma if untreated.



Always test for ketones if a person with type 1 or type 2 diabetes presents to general practice and is unwell (see page 46 for ketone levels)

When a person with diabetes is acutely unwell, the role of primary care is to ensure that person is

already well educated about sick day management well supported with their diabetes management

while treating the underlying cause of the illness

Type 1 Diabetes

- patients can develop diabetic ketoacidosis (DKA) when unwell with an increase in levels of counter regulatory hormones which leads to hyperglycaemia and ketone bodies due to not enough insulin present
- They will often require a temporary increase in insulin which can include basal rates and supplementary rapid acting insulin 1-2 hourly if ketones are present and while they remain unwell.
- Dehydration must be avoided with frequent fluid replacement hourly (aim for at least a cup or glass hourly).
- Patient to contact their Diabetes nurse specialist or present to ED if hyperglycaemia/ ketones/vomiting/abdominal pain/fruity breath or shortness of breath are present

Almost all sick day management can happen in primary health care setting.

Preparing the patient prior to them getting sick is important.

On the following 2 pages is a sick day leaflet for patients with T2DM that you can copy for your patients or download at https://www.pinnaclepractices.co.nz/resources/diabetes-sick-day-advice/

Sick day management for people with type two diabetes.

When you are sick you may notice that your blood glucose levels change. This can be because your appetite changes, but also because infections, trauma, stress and other illnesses can cause your blood glucose levels to go up or down.

It can be difficult to manage your diabetes on your own when you feel crook, so it is important to let a whānau member, friend or neighbour know when you are unwell.



Drink:

If you are vomiting and not eating or drinking enough, you can quickly become dehydrated. Sip at least one glass of sugar free drink every hour. (Water, sugar free soft drink, tomato juice, tea, weak coffee)







Food:

If you are unable to eat your usual meals, something small and full of energy each hour can help. Some examples are orange juice, milk, soup and toast, fruit yoghurt, 4 – 6 water crackers, 1 small banana

Checking:



Check your blood glucose levels often – every 2 or 4 hours if you are really sick. More often if needed. This is so you can react if your blood glucose is too high or too low. You might need to call your nurse for help, and she will ask how your blood glucose levels have been.

Healthline is a free 24-hour telephone health advice service. Phone 0800 611 116 for any health information.



Medicine:

If you are feeling a little unwell, its really important that you still take the medicines the doctor has prescribed for you.



If you are really sick and if your glucose levels are higher than normal – contact your nurse for help. You may neeed to change the medicine you take.

If your glucose levels are lower than normal – you may need less of your insulin or tablets that you take.



Hypo – if your blood glocose is too low or under 4 mmols.

If your blood glucose levels are under 4 mmols – your need to treat this 'hypo' with a good dose of sugary drink or food – a glass of regular fizzy or orange juice, 6 teaspoons of sugar in warm water or 18 jelly beans. Test your blood glucose levels again in 15 minutes, if you are still under 4 mmols, treat again with the sugary drink or jelly beans, but if you have blood glucose above 4 – have something to eat like your next meal, a banana or a slice of toast.



Phone your doctor or nurse if you

Cannot keep your blood glucose above 4 mmol
Are too sick and have no support person
Are passing lots of urine or have abdominal pain
Have nausea, vomiting or diarrhoea for more than 12 hours
Feel drowsy, weak or confused
Have chest pain
Have difficulty breathing

Healthline is a free 24 hour telephone health advice service.

Phone 0800 611 116 for any health information

ORAL HEALTH

People with diabetes (type 1 & 2) have a high prevalence of oral problems such as dental caries, xerostomia (reduced or absent saliva flow causing dry mouth), periodontal disease, taste problems and oral infections. Periodontal disease which is an inflammatory condition destroys the connective tissue surrounding the teeth which can lead to tooth loss. These conditions are caused by poorly controlled blood glucose levels which cause susceptibility to bacterial infections in the mouth.

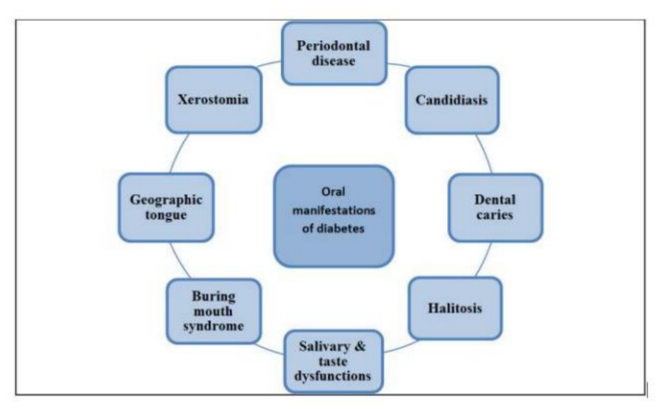


Figure 5: Oral manifestations among diabetic patients [23] [41] Open Access Macedonian Journal of Medical Sciences. 2018.

Patient education includes being aware of the risk for periodontal disease and the importance of maintaining oral health as part of their diabetes management. This includes regular dental checks and care, with collaboration between patient and all health care professionals. Advice should include tooth brushing twice daily, with dental floss once a day to control plaque. The use of mouthwashes to prevent caries and plaque build-up may be useful for prevention of periodontal disease.

PLANNED/ELECTIVE SURGERY

Diabetes Management for Gastroenterology Procedures

On most occasions these are outpatient procedures. All situations should be discussed with GP and practice nurse. However, the Endoscopy Unit have specific endoscopy protocols for the management of patients with diabetes who are undergoing endoscopic procedures.

These include upper endoscopy, endoscopic ultrasound, endoscopic retrograde cholangiopancreatography, bronchoscopy and anterograde double balloon.

There is also the preparation for lower endoscopic procedures which include colonoscopy and retrograde double balloon procedure, Klean-Prep or Pico-Salax preparation and capsule endoscopy. These can either be morning or afternoon procedures but for people with diabetes we would expect that these procedures are scheduled for the mornings.

Patients will need an **insulin reduction** for all these procedures, particularly if they need to be nil by mouth for an extended time or require bowel prep prior to surgery. This will involve the days prior to the procedure and the day of procedure, and the patient's current blood glucose profile must also be considered. This is particularly relevant if they have tight control or you feel that their current dosage might be a bit too high. It will also depend on whether they are using basal/bolus or premixed insulins.

It may be necessary to stop or reduce other medications such as Metformin or Sulfonylurea prior to the procedure. For procedures involving bowel prep, Empagliflozin (Jardiance) should be stopped 3 days prior and the day of (4 days in total) to reduce the risk of diabetes ketoacidosis (DKA).

Other Elective Surgery

If your patient is referred for an elective procedure, it is imperative to support good glycaemic control.

- Stop Metformin and Empagliflozin (Jardiance) 2 days prior to planned surgery
- Patients with HbA1c ≥70 mmol/l may be put "on hold" until their diabetes is better managed. Higher HbA1c indicates
 - Twice as likely to get surgical site infection
 - † risk post-operative myocardial infarction
 - post-operative renal failure (AKI)
- This means at least another 4 months wait for their operation

Plan ahead and maximise glycaemic control as soon as the patient is referred for surgery

TRAVEL GUIDELINES

There are a number of recommendations to be given to people with diabetes so that they can travel safely. These include planning travel with the below safeguards in mind:

- Have a medical consultation prior to travel to assess diabetes control and check immunization status
- Ask GP to provide a covering letter for travel stating that the person has diabetes requiring use
 of insulin, syringes/pen devices, monitoring equipment such as insulin pumps/continuous
 glucose monitoring devices
- Check with airline if continuous monitoring devices can be operated during flight
- Advise to use travel insurance
- Advise to find out from airline about security guidelines in countries being visited
- Carry supplies on board with the person in their hand luggage
- Do not put insulin supplies in checked in luggage as it will be exposed to extreme temperatures which may degrade the insulin (lose efficacy)
- It is a good idea to also carry a printed list of prescriptions in case of medication loss or theft of bag.



If a person is on insulin and changing time zones, they will need to consider how they will change their insulin to meet their needs while away. Any advice given will depend on if the patient is on basal only or basal plus rapid acting insulins. Work out when the most likely time is that the person will be having a meal, and how they can change into the new time zone. Remember there is quite a difference in the insulin needs between a person with type 1 diabetes (contact the diabetes clinic for advice) with absolute insulin deficiency and a person with Type 2 diabetes and some residual beta cell function.

MEDICATION USE REVIEW

A 'Medicines Use Review' aims to improve understanding and adherence to medicines by identifying and addressing factors linked to non-adherence behaviours.

Patients (and their family/whanau) with complex difficulties in understanding and adhering to medications, receive personalised education and support to improve self-management.

The Pinnacle extended care team pharmacist is available for medicine use review. Send an BPAC stating medication management support.



PRE-PREGNANCY PLANNING

It is important that a woman starts the pregnancy with well controlled diabetes so that the baby's growth and development is optimised.

Questions regarding contraception and pregnancy planning are embedded on the common form as prompts as 50% of pregnancies are not planned.

Is there a potential to get pregnant?	O Yes	O No	Con	sider pre	conception advice where appropriate
Do you need to discuss contraception	w/patient?	OY	es	O No	
Patient's form of contraception?					

Women with pre-existing diabetes who are planning a pregnancy should be supported to achieve excellent glycaemic control prior to conception. This means blood glucose of 4 − 8mmol/L, or an HbA1c ≤48mmol/mol without hypoglycaemia. They will require an appointment with their prescriber to maximise therapies and start folic acid.

If HbA1c is elevated, ensure contraception is available or prescribed until glycaemic control is achieved.

Contraindicated in pregnancy:

- Statin
- ACE
- SGLT2i, GLP1RA, DPP4
- Sulphonylurea—consider starting insulin prior to pregnancy.

Recommended prior to pregnancy:

- Folic acid 5mg (not the standard 0.8mg)
- Retinal screening within last 6 months
- Stop smoking and alcohol
- Weight management

The aim for pregnancy is a healthy baby and a well mother. By focusing on pre pregnancy care, the whānau start the pregnancy with the best outcomes in mind. For 3 months prior and during pregnancy, aim to achieve normoglycemia so that the baby is conceived and develops in conditions as close to a pregnancy uncomplicated by diabetes as possible.

At times, this means a lot more effort from the woman with food choices, activity and checking blood glucose. Recommended reading: https://www.nice.org.uk/guidance/ng3/chapter/1-recommendations#preconception-planning-and-care-2

Te Whatu Ora - Taranaki has a pre-conception pregnancy planning service – referrals via BPAC.

Refer all women with pre-existing diabetes who become pregnant to the team, referrals can also be initiated by an LMC

PAEDIATRICS AND YOUTH

Referrals can be made to the Te Whatu Ora – Taranaki paediatric team for children up to the age of 15. Contact Clinical Nurse Specialist Gina Newman via email or phone

If a child presents with a concurrent illness, the role of general practice is to assess and treat the illness with the support of the paediatric team as required.

Paediatric.DiabetesCNS@tdhb.org.nz or on 027 5577 110

For rangatahi aged 15-25, the youth diabetes service at Te Whatu Ora – Taranaki is run by nurse practitioner Dorothy Larsen dorothy.larsen@tdhb.org.nz or 027 801 0065 who accepts referrals for both type 1 and 2 diabetes in this age range.

Taken from the Starship website for parents: What to do when your child is sick: https://www.starship.org.nz/sick-days

Children and young people with diabetes generally don't get sick any more often than other children if their diabetes is well managed. However, illness can have a significant impact on diabetes. The stress hormones produced during illness can cause changes to blood glucose levels. Levels can go high or low depending on the type of illness. Infections that cause fever and pain often cause high blood glucose levels. Gastroenteritis (vomiting and diarrhoea) can cause low blood glucose levels. Vomiting can also be a sign of not enough insulin.

If your child is sick:

- Always give insulin, but call the diabetes doctor/nurse for advice on changes to the usual amount
- Take your child to your GP for assessment and treatment of the underlying illness
- Test blood glucose levels 2 hourly
- Check blood or urine ketones 2 hourly (see page 46 for results and what to do)
- Encourage your child to drink plenty of fluids

If your child is unable to eat:

- Always refer to the child's sick day management plan from the paediatric team
- If blood glucose levels are under 10, give fluids with sugar such as diluted juice or jelly (not sugar free)
- If blood glucose levels are 10 or higher, give water or sugar-free fluids
- If your child has been vomiting or has diarrhoea, Pedialyte™ may be recommended by the diabetes doctor or your GP (this can be purchased from the pharmacy)
- Relieve the symptoms of pain and fever with Paracetamol or Ibuprofen

Even teenagers who usually manage their diabetes independently need to be looked after when they are sick .

APPENDICIES

Appendix 1: Diabetes Model of Care in Tarana	ki & General Practice Error!
Bookmark not defined.7	
Appendix 2: Key Contacts – Extended Care Tea	am Error! Bookmark not
defined.8	
Appendix 3: Key Contacts – Industry Reps8E	rror! Bookmark not defined.
Appendix 4: NZSSD T2DM Guideline Alogorithr	m Error! Bookmark not
defined.1	
Appendix 5: NZSSD Insulin Algorithm Err	or! Bookmark not defined.2
Appendix 6: Key Words Err	ror! Bookmark not defined.3
Appendix 7: Recalls Err	or! Bookmark not defined.4

Appendix 1: Diabetes Model of Care in Taranaki & General Practice

Diabetes Integrated team (DIT)

NORTH TARANAKI ONE

NORTH TARANAKI TWO

CENTRAL / SOUTH TARANAKI

- 1. City West Medical Centre
- 2. Family Health Centre
- 3. Tukapa Medical Centre
- 4. Dr Bellomo
- 5. Healthspace
- 6. Medicross GP Clinic
- 7. Vivian Medical Centre
- 8. Care First Group
 - Westown MC
 - · Merrilands MC
 - Moturoa MC
 - Bell Block MC

- 1. Devon Medical Centre
- 2. Tui Ora Family Health
 - Whaitara HC
 - Westown HC
- 3. The Nest
- 4. Eastside Medical Centre
- 5. Full Circle Medical Centre
- 6. PHCL Group
 - Moa Medical Centre
 - Central Medical Centre
- 7. Omni Group
 - Parklands MC
 - Strandon Health MC

- 1. Mountainview Medical Centre
- 2. Ngati Ruanui Health Care
 - Hawera HC
 - Patea HC
- 3. Dr Blayney*
- 4. South Taranaki Rural Health*
- 5. Ngaruahine Iwi Health Centre
- 6. Coastal Group
 - Opunake MC
 - Avon MC
 - Oakura MC
 - Waiwhakaiho MC
- 7. Eltham Health Centre
- 8. Regan St Health Centre

**Dedicated cluster relationship Clinical Specialist - Lauren + CNS **Dedicated cluster relationship Clinical Specialist - Lauren + CNS

**Dedicated cluster relationship Clinical Specialist - Lauren + CNS

Kaitautoko Mate Huka Tui Ora / Ngaruahine / Pinnacle MHN / Ngati Ruanui

Consultant Physician GPSI Diabetes, Specialist Dietitian, Endocrinologist,
Psychology & Foot Protection Service

All Clinical roles are diabetes specialist

V8 July 2024

* Not in PHO

** Dedicated Cluster relationships - Enhancement & Capability Support Provide support, education and training for both health professionals and patients.











Appendix 2: Key Contacts – Extended Care Team

https://www.pinnaclepractices.co.nz/programmes/extended-care-team-referral-service-taranaki/

The extended care team is available to compliment the care you provide to your patients.

The team includes a number of disciplines both clinical and non-clinical who are able to support your patients with the complexities of their long-term condition management.

The care provided is community-based and may be interdisciplinary and comprise a combination of group, peer and one on one support

This care is provided across the region.

All referrals are triaged by the team and the result of triage will be communicated back to you via BPAC. To be eligible for these services patients must be eligible for publicly funded care in New Zealand.

Priority will be given to Māori/Pacific patients

Eligible patients are those who are:

- Diagnosed with a long-term condition e.g., Type 2Diabetes, CHF, CVD, COPD or identified as high risk of developing a long-term condition
 - Aged over 15 years
 - Not currently under care of secondary care allied health

And have

- Identified concerns/issues regarding medication management
- Identified concerns/issues regarding nutrition
- Difficulties self-caring/self-managing/accessing traditional GP services
- Identified concerns/issues regarding lifestyle/environmental support
- Identified concerns/issues regarding communication/memory/disengagement with health services

The team consists of dietitians, pharmacists, social workers, health improvement practitioners and Kaitautoko Mate Huka.

Referral is via BPAC and to the team, not to individual clinicians (ie, not specifically to a dietitian, or a pharmacist). To allow for successful triage, please make sure it includes

- The main reason for your referral in the Clinical Details page of the BPAC EReferral this is a mandatory field.
- Any relevant test results
- Any relevant goals/care plan information
- Any other known support agencies/services involved

Appendix 3: Key Contacts – Industry Reps

Free phone for all New Zealand 0800 283684

Sanofi New Zealand

T: +64 9 580 1810 (option 1) F: +64 9 580 1811
Level 8, James and Wells Tower, 56 Cawley Street, Ellerslie, Auckland 1051, New Zealand
PO Box 12851 Penrose, Auckland, New Zealand.









Novo Nordisk

Hamza Rikabi Diabetes Care Specialist **0274445722** HARB@novonordisk.com

Customer Careline please call on 0800 733 737



Olivia Solomons PH: 021 1959634

olivia.solomons@pharmaco.co.nz







THERESA DAVIES

(She/her) TERRITORY MANAGER

027 278 6671 | 0800 106 100 | +64 9 414 0318

www.mediray.co.nz

Nicola Gibson-Groot: 0274993101

nicola.gibson-groot@boehringer-ingelheim.com







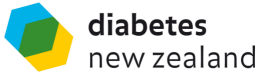






For patients to contact:

URL www.diabetes.org.nz Email info@diabetes.org.nz Phone 0800 342 238



MANAGEMENT ALGORITHM FOR TYPE 2 DIABETES



INITIAL MANAGEMENT

Confirm the diagnosis and type of diabetes Determine individualised glycaemic target

Lifestyle management

Education, support, healthy eating + exercise Essential at all times throughout duration of diabetes

Start unless contraindicated Increase to maximal tolerated dose or 2 g per day

Weight management

- Set individualised weight management plan if overweight or obese
 5% sustained total body weight loss is associated with improvement in metabolic parameters 10–15% sustained total body weight loss is typically needed for remission of diabetes

The target HbA1c for most patients with type 2 diabetes is < 53 mmol/mol

- If HbA1c > 64 mmol/mol at diagnosis consider starting additional agent with lifestyle management and Metformin to reach target
 - If cardiovascular and/or renal disease and/or heart failure preferably SGLT2i or GLP1RA (see below)
 - If no cardiovascular or renal disease and no heart failure preferably DPPIVi
- · Consider starting insulin therapy immediately if:
 - Symptoms of hyperglycaemia/insulin deficiency and/or HbA1c > 90 mmol/mol Suspicion of type 1 diabetes or loss of pancreatic function

Risk of

hypoglycaemia

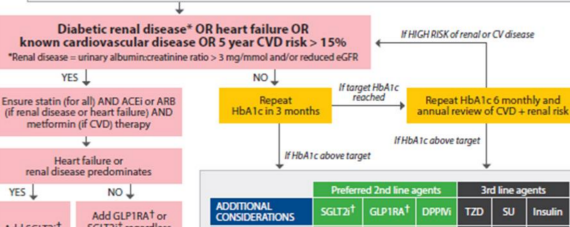
Mean

in HbA1c (mmol/mol)

cardiorenal benefits

Effect on weight

Funded



Rare

SA only^T

Rare

15

Yes

SA only[†]

Rare

5 - 10

No

Rare

15

Yes

٠

Yes

15

No

٠

Yes

Any

No

٠

SGLT2i[†] regardless of HbA1c if no Add SGLT2i[†] regardless contraindications. of HbA1c GLP1RA likely if no contrapreferable if indications cerebrovascular (HbA1c needs to be >53 mmol/mol for funding) disease predominates (HbA1c needs to be >53 mmol/mol for funding) If unable to tolerate or HbA1c remains

above target

GLP1RA[†] preferred next therapy after SGLT2i[†] SGLT2i[†] preferred next therapy after GLP1RA[†] (dual SGLT2i/GLP1RA therapy is not currently funded)

Alternative agents include: DPPIVi if not on GLP1RA Thiazolidinediones (TZD) if no heart failure Sulfonylureas (SU) Insulin

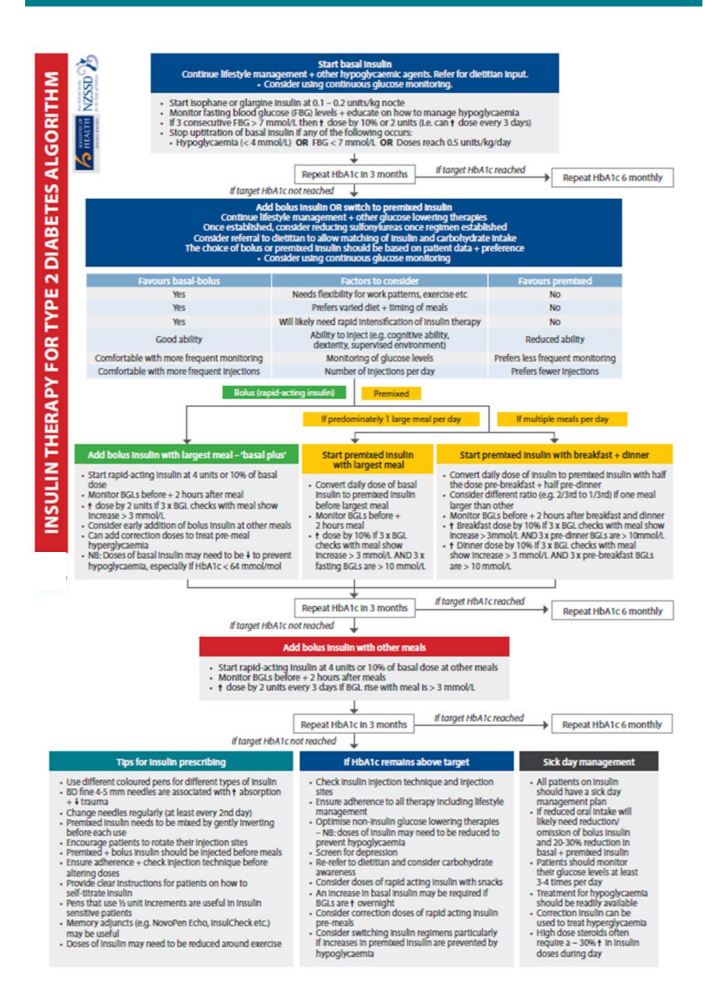
†SA criteria for SGLT2I and GLP1RA (all regulred and same for both classes)

- Patient has type 2 diabetes with an HbA1c
 53 mmol/mol despite > 3 months of regular use of at least one glucose lowering therapy (includes metformin)
- · The patient is of Māori and/or any Pacific ethnicity OR has known diabetic renal disease OR known CVD OR 5 year CVD risk > 15% OR a high lifetime CVD risk due to onset of diabetes during childhood or as a
- · The patient is not on funded SGLT2i and GLP1RA therapy at the same time

Escalate therapy + repeat HbA1c every 3 months until target reached

- · May require multiple agents including insulin therapy
- · Ensure adherence to lifestyle management + medications
- · Re-refer for dietitian input if appropriate
- Repeat HbA1c 6 monthly once target reached
- Assess CVD and renal risk at least annually
- · Continue standard care to reduce CVD risk e.g. statins, antihypertensives (esp. ACEi in diabetic renal disease) etc.

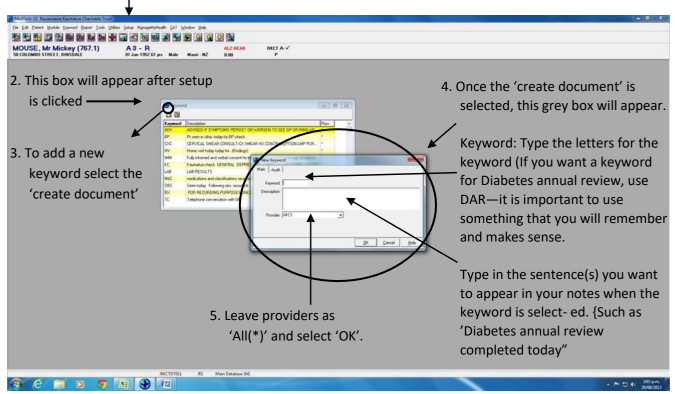
Appendix 6: NZSSD Insulin Algorithm



Appendix 7: Key Words

Using **KEYWORDS** can make writing notes easier and helps keep documentation consistent between users. This is an example on MedTech—your PMS system will have similar capabilities. Talk with your administrator to set up keywords for your practice:

1. After opening MedTech locate the option for:



Now to practice:

Using MedTech, open the Mickey Mouse practice screen.

In the clinical notes type. DAR (full stop DAR) and press 'Enter', and it will automatically self-populate what you set up.

The headings listed here guide what you are going to record. You start typing after each heading and then press **ENTER**. This will move to the next heading down the consult and you can start typing beside it, repeating until all headings have an entry beside them:

E.g. Diabetes annual review completed today

PLAN: HbA1c remains high, requires more oral medication—task to GP.

CLINICAL FINDINGS: BP LIPIDS WEIGHT all stable

<u>NUTRITION:</u> Food plan completed 2016—discussed today and remains very similar. Portion size the biggest issues along with eating out at weekends.

EXERCISE: sedentary at office but has been walking across carpark and up 4 sets stairs.

Try to use keywords for longer blocks of typing.

Aim: to have keywords you can recall easily and that will help save you time.

Appendix 8: Recalls

Every practice will have a different process/system - the key, is to <u>have</u> a system and a key person assigned to:

- monitor/do the recalls
- Be familiar with the recall list and BPI information

Below is a mock-up recall letter - you will need to adapt this for individual needs (annual, 3-month monitoring appointment, Type 1 diabetes, Type 2 diabetes).

The majority of clinics are contacting patients with verbal and paper invitations and we know that those who do not respond to the initial letter or the five phone calls are most probably the people who would benefit the most.



DAR Recall Process

A designated person initiates DAR recalls at the beginning of each Quality quarter (July, October, January and April). If there is a query, this person can clarify recalls. The essence is that no person misses their opportunity to be seen and discuss their diabetes management goals and aspirations.

- 1. Print off the BPI list of High Needs "Not Achieved" Patients with HbA1c >64mmol/mol Refer to this list when sending DAR invites.
- 2. Create the Recall Contact List for all patients requiring DAR during the next Quarter, using the toolbar icon or via the "Module" tab.
- 3. For each patient on the Recall Contact List:
 - Check Classifications: If Type 1 Diabetes, check if patient is under secondary review, and if so, send a task to the designated Diabetes Nurse to decide if a Virtual DAR is best.
 - Check screening: Has DAR already been completed? If so, check if Recall is correct
 - Check Outbox: Has DAR invite/lab form already been sent?
 - Check Inbox: Have lab tests already been completed?
 - Check Daily Record, Classifications and Tasks for any relevant information/exceptions to DAR.

Exceptions to DAR Recall:

For any of the following, send a Task to the designated Nurse to advise DAR is due. The Nurse can liaise as appropriate with either the Doctor or other service provider.

- Patients who are in Rest Home care (See below)
- Patients who are palliative
- Patients who are being intensively managed by a designated Diabetes Nurse
- Patients who are under the care of Specialists services

First Contact (Letter)

- Print DIABAR letter (DAR invite)
- Print DAR lab request. Check Tasks for any other lab tests which are also due (Liaise with Doctor as needed)
- Forward letter and lab form together to doctor for signing, before being mailed
- If the patient is identified as "High Needs" on the BPI list: Send a task to the designated nurse to follow up in 2 weeks.
- Do not reschedule recall at this stage, to keep the Patient Prompt Recall active
- Adjust "R#" in the Recall Contact List to "1", to indicate 1st contact made

Second contact (phone)

After 2 weeks if "High Needs" patients, and 6-8 weeks if standard DAR

- Check Screening: Has DAR been completed, and Recalls updated?
- Check Appointments: Has DAR been booked?
- Check Daily Record, Classifications and Tasks for any new relevant information/exceptions before phoning patient
- Check Inbox: Have lab tests been completed?
- \circ \Rightarrow If not, phone patients to check letter and lab-form were received, and to remind them to complete lab tests and book DAR appointment
- ⇒ If lab tests have been completed, phone patient to book DAR appointment. Consider offering a doctor's appointment if prescription or other review is due. The DAR appointment should be scheduled before the Doctor's appointment.
- Do not reschedule recall
- Adjust "R#" in the Recall Contact List to "2", to indicate 2nd contact made.

As with all patient phone contact, check contact details are correct, check and advise if other recalls are due (e.g., cervical smear) and update smoking status as appropriate.

Third Contact (letter)

After 2 weeks if "High Needs" patients, and 6-8 weeks if standard DAR

- Check as per Second Contact above
- If no appointment made send DIABR2 letter with a copy of lab test request if not yet completed
- Adjust "R#" in the Recall Contact List to "3", to indicate 3rd contact made

If DAR incomplete after a further interval, send a Task to GP/designated Diabetes

Nurse for review.

Rest Home DARs

Rest Home Contact Information:

Rest Home DAR Process

- If all people with diabetes within the rest home come to your clinic,
- The Diabetes Nurse(s) or HCA responsible for the Rest Home DARs will coordinate these DARs, following the standard DAR Recall Process checks
- A single covering letter is sent to the Nurse in Charge/Manager at the Rest Home, with all the DAR lab request forms for the residents due to their DAR.
- Forward letter and lab forms together to doctor for signing, before being sent to the Rest Home
- Monitor patients on task list. Make notes of the dates lab request form were sent, when results are received, and leave on task list until DAR documentation complete
- When results are back, arrange visit with the Rest Home Nurse in Charge/Manager
- On the d ay of t h e Rest Ho me visit:
- Check Rest Home has a list of patients being seen, and request patient file be available to include copy of medication chart, current weight and height.
- Take required equipment for DAR, including sphygmomanometer, foot check monofilament, tape measure, steri-gel, as well as DAR forms, and laptop if available
- Invoice via reception on the day seen.
- Complete PMS notes and submit Common Form. Task can then be completed
- Check and adjust Recalls as appropriate
- Send task to doctor advising DAR has been done

Diabetes Annual Review Decline Process

- If a Patient declines to have a DAR, document this in their notes using a keyword you have set up adjusting as appropriate:
 - "Face to face/phone discussion re diabetes annual review invitation. Pt declines to have DAR her as: seen elsewhere .../ declines to participate. DIAP code entered as incomplete, with declined, added to note line"
- In "Screening" under "Outcome" select "Incomplete Screening (IN). Type "declined" in the 'Note' field. This enables available data to still be added.
- Reschedule Recall for 1 year and send a task to the designated nurse to advise of the decline
- Designated Diabetes Nurse to complete Screening template as much as possible, as for "Virtual DAR"

Diabetes Monitoring Recall Guideline

Diabetes Monitoring Recalls are completed by the designated Diabetes Nurse between patient DARs - as a means of providing ongoing Diabetes support to the patient and based on goals set at DAR.

To create the recall contact List:

Process:

- When the DAR is completed, a recall for "Diabetes Monitoring" (DCIP) is automatically generated for 3-6 months' time
- Monitoring notes can be added into DCIP (e.g., repeat HbA1c and BP in 3 months). As a minimum, all patients with diabetes should have their HbA1c checked 6-monthly.
- The designated Diabetes Nurse is responsible for implementing the Diabetes Monitoring
 Process for their own patients. This may include reminding patients to complete lab tests,
 reviewing current Diabetes control and management, medications reviews and liaising with the
 Doctor as required

Before contacting patients, the designated Diabetes Nurse should check the patient records for any new information, including the Daily Record and Inbox.

- Make an entry in the notes using the Diabetes Monitoring classification, noting current Diabetes management and control, any recent investigations, and advice given including intensification of treatment
- Liaise with/send task to GP if further input required or send task to self to provide further follow- up.
- Reset Diabetes Monitoring Recall as appropriate.

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