

Taddle Creek

Family Health Team

Title: Requisition of Laboratory Investigations **Number:** TCFHT- and Prescribing Diabetes Supplies for the MD16

Management of Diabetes or Prediabetes

 Activation Date:
 09-Sep-2014
 Review
 12-Jun

 Date:
 2021

Next Review Date: 12-Jun-2022

Note: Jun 2016 review resulted in a change; ability to order non-fasting lipid profile. Change approved at Jun 14-16 Board Mtg (see minutes) thus negating necessity to get authorizers to re-sign.

Sponsoring/Contact Person(s)

(name, position, contact particulars):

Dr. Mitch Vainberg

790 Bay Street. Suite 300 Toronto, Ontario M5G 1N8

416-960-1366

Karen Finch, Registered Nurse, Certified Diabetes Educator

790 Bay St. Suite 508Toronto, Ontario M5G 1N8 **416- 204-1256, x502**

Sherry Kennedy, Executive Director – skennedy@tcfht.on.ca

790 Bay Street, Suite 306, Toronto, Ontario M5G 1N8

416-260-1315, x307

Order and/or Delegated Procedure: Appendix Attached: No X Yes

Title: Appendix C – Performed Controlled Acts and Procedures (CAPs) Implemented Under this Directive

The implementers are authorized to perform the following activities, in accordance with Appendix C when all conditions in this directive and the attached appendices are met:

- 1) Requisitioning of Laboratory Investigations, by implementers, for patients of the Taddle Creek Family Health Team (TCFHT) Primary Health Care Providers (PCPs) or of Consulting Endocrinologist and who meet specific indications described within this directive.
- 2) Prescribe Diabetes Supplies (flash glucose monitor, flash glucose sensors, glucose meter, glucose meter strips, blood ketone test strips, needles for insulin pens and lancets)

Recipient Patients: Appendix Attached: __ No X Yes

Title: Appendix A – Authorizer Approval Form

Recipients must:

- Be active patients of a TCFHT primary care provider or Consulting Endocrinologist who has approved this directive by signing the Authorizer Approval Form
- Have a diagnosis of Diabetes Mellitus (type 1 or 2) or Prediabetes
- Meet the conditions identified in this directive

Authorized Implementers:

Appendix Attached: ___ No _X_ Yes

Title: Appendix B – Implementer Approval Form

Implementers must be TCFHT employed Regulated Health Care Providers or Physician Assistant (under the supervision of a physician).

Implementers must complete the following preparation and sign the Implementer Approval Form:

- Must be Certified Diabetes Educators (CDEs) who practice according to the most current recommendations for the management of diabetes
- Practice according to Diabetes Canada's (DC's) most current Clinical Practice Guidelines
- Assess their own knowledge, skill, and judgment to competently perform these directives
- Must complete the Implementer Competency Checklist for Prescribing Diabetes Supplies (Appendix E) prior to signing the Implementer Approval Form

Appendix Attached: ___ No _X Yes

Title: Appendix C – Performed Controlled Acts and Procedures (CAPs) implemented under this directive

Indications:

- Each action/procedure under this directive will be implemented in the context of the existing
 PCP or Consulting Endocrinologist-patient relationship and as part of the medical diagnosis and
 plan of care established by the PCP. These actions/procedures will be implemented without
 specific prior discussion (but as part of the plan of care) as per the indications and
 contraindications for each of the directives.
- Specific indications for each laboratory investigation ordered under this medical directive can be found in Appendix C
- Specific indications for flash glucose monitoring, blood glucose monitoring, blood ketone testing and use of needle tips can be found in Appendix C

Contraindications:

Indications described in Appendix C are not met

Consent: Appendix Attached: X No Yes Title:

- Patient's consent is implied for implementer to provide lab requisition or provide prescriptions
 for flash glucose monitoring, blood glucose monitoring, blood ketone testing or needle tips, as
 patient has presented seeking support with diabetes management, and is a Family Health Team
 patient or patient of the Consulting Endocrinologist, where interprofessional practice is
 expected
- Patient informed of purpose of testing, including when results will be available and contact information to review results (if not contacted by PCP)

Guidelines for Implementing the Order/Procedure:

Appendix Attached: No X Yes

Title: Appendix C – Performed Controlled Acts and Procedures (CAPs) implemented under this directive Appendix D – Sample Lab Requisition

Requisitioning of Laboratory Investigations

- 1) Identify need for laboratory investigation (blood work) and determine whether indications described in Appendix C are met.
- 2) Ensure that no recent blood work has been undertaken that would result in duplication of testing.
- 3) Explain the purpose of the test to the patient
- 4) Generate a laboratory requisition using the supervising PCP/Authorizers initials.
- 5) Lab Requisition should be signed as below:
 - Signature
 - o Implementer Name/Primary Care Provider Name (Medical Directive TCFHT-MD16)
- 6) Send a message in Practice Solutions to the PCP indicating that a lab requisition has been provided.
- 7) PCP will receive completed lab requisitions and forward them to implementers as needed e.g. if earlier follow up with implementer is required
- 8) Implementer documents that the requisition was provided and follow up plan in the eMR.

Prescribe Diabetes Supplies

- 1) Identify need for prescribing diabetes supplies and determine whether indications described in Appendix C are met.
- 2) Patient education provided on self-monitoring of blood glucose, flash glucose and/or blood ketones as per DC's most current Clinical Practice Guidelines
- 3) Enter a Rx for diabetes supplies following the steps outlined in DEP 12 Prescribing Diabetes Supplies and/or making Medication Changes in Practice Solutions (Program Folders/Diabetes/Procedures/DEP 12 Prescribing Diabetes Supplies and/or making medication changes in Practice Solutions)

Documentation and Communication: Appendix Attached: __ No __ Yes Title:

- Documentation in the patient's eMR needs to include: name and number of the directive and name of the implementer (including credential)
- Information regarding implementation of the procedure, the patient's response and follow up plan should be documented in the patient's eMR, in accordance with standard documentation practices (College of Nurses, 2008).

Review and Quality Monitoring Guidelines:	Appendix Attached: _	No	_ Yes
Title:			

- Review will occur annually. Review will involve a collaboration between the authorizing primary care providers and the approved implementers.
- If new information becomes available between routine reviews, such as the publishing of new
 clinical practice guidelines, and particularly if this new information has implications for
 unexpected outcomes, the directive will be reviewed by an authorizing primary care provider
 and a minimum of one implementer.
- At any such time that issues related to the use of this directive are identified, TCFHT must act
 upon the concerns and immediately undertake a review of the directive by the authorizing
 primary care providers and the authorized implementers.
- This medical directive can be placed on hold if routine review processes are not completed, or
 if indicated for an ad hoc review. During the hold, implementers cannot perform the
 procedures under authority of the directive and must obtain direct, patient-specific orders for
 the procedure until it is renewed.

References

Diabetes Canada. (2018). Clinical Practice Guidelines for the Prevention and Management of Diabetes in Canada.

College of Nurses of Ontario. (2008). *Practice Standard: Documentation*. Retrieved from https://www.cno.org/globalassets/docs/prac/41001 documentation.pdf

Pearson et al. (2021). 2021 Canadian Cardiovascular Society Guidelines for the Management of Dyslipidemia for the Prevention of Cardiovascular Disease in the Adult Canadian Journal of Cardiology 2021 Mar 26;S0828-282X(21)00165-3. doi: 10.1016/j.cjca.2021.03.016. Online ahead of print. Retrieved from: https://www.onlinecjc.ca/article/S0828-282X(21)00165-3/fulltext#seccesectitle0037

Goldenberg, R.M., Cheng A.Y.Y, Punthakee, Z., et al. 2011. Use of glycated hemoglobin (A1C) in the diagnoses of type 2 diabetes in adults. *Canadian Journal of Diabetes*; 35: 247-248.

Merck Sante. Product Monograph: Glucophage.<u>http://products.sanofi.ca/en/glucophage.pdf</u> Version 5.0

Canadian Insulin Injection Recommendations: FIT Technique Plus, 2020. Retrieved from http://www.fit4diabetes.com/canada-english/fit-technique-plus/

Fit Forum for Injection Technique Canada. (2020). Fit Forum for Injection Technique Canada: Recommendations for Best Practice in Injection Technique. Retrieved from http://www.fit4diabetes.com/files/7816/0803/3133/FIT_Recommendations_2020.pdf

Note: This medical directive is for the routine monitoring of laboratory investigations for the management of diabetes or prediabetes and does not include other laboratory investigations (ALT, AST, CK or CPK, CBC etc.), which are recommended for starting or monitoring the effects of medications e.g. oral antihyperglycemic medications, statin medications etc.

Appendix A:

Authorizer Approval Form

Name	Signature	Date
	 	_ '
	 	-

Appendix B:

Implementer Approval Form

To be signed when the implementer has completed the required preparation, and feel they have the
knowledge, skill, and judgement to competently carry out the actions outlined in this directive.

Name	Signature	Date
	-	
		<u> </u>

Appendix C

Table 1: Controlled Acts and Procedures (CAPs) Implemented Under this Directive

Laboratory Investigation	Indications
Fasting Blood Glucose (FBG) & Glycated Hemoglobin (HbA1C)	Every 3 months when glycemic targets are not being met and/or when diabetes therapy is being adjusted. Every 6 months should be performed in adults during periods of treatment and lifestyle stability when glycemic targets have been consistently achieved. Every 6-12 months is recommended for people with prediabetes. FBG should be obtained after an 8-12hr fast. A Random Blood Glucose (RBG) along with an HbA1C should be considered for patients at high risk for hypoglycemia e.g. those taking insulin, frail elderly etc. An HbA1C may be misleading in some people with various hemoglobinopathies, iron deficiency, hemolytic anemias, and severe hepatic and renal disease. A fructosamine test can be used in these cases for a cost of approximately \$35. The RN or RD to consult with the PCP and can obtain a verbal order for this test if indicated.

Lipid Panel (total cholesterol, triglycerides, HDL – cholesterol, LDL-cholesterol, total cholesterol: HDL-C ratio)

A lipid profile (TC, HDL-C, TG, and calculated LDL-C) should be measured at the time of diagnosis of diabetes. If lipid-lowering treatment is not initiated, repeat testing is recommended yearly. More frequent testing (every 3-6 months) should be performed after treatment for dyslipidemia is initiated (lifestyle and/or medications).

A non-fasting lipid profile should be considered for most patients to improve adherence and to lower the risk for hypoglycemia. New evidence indicates minimal differences exist between fasting and non-fasting HDL, LDL, and total cholesterol levels. The differences that occur are less than the within-person variability from repeat lipid testing. Tests of non-fasting HDL and non-HDL levels correlate with future CVD events. Although triglycerides are most susceptible to change without fasting, triglycerides contribute minimally to total cholesterol levels, and triglyceride levels are not consistently associated with CVD.

CCS 2021 guidelines indicate that for any patient with triglycerides > 1.5 mmol/L, non-HDL-C or ApoB be used instead of LDL-C as the preferred lipid parameter for screening at a cost of \$25. RN or RD to consult with the PCP and can obtain a verbal order for ApoB if indicated.

A fasting lipoprotein profile should be obtained after a 10hr-12hr fast, preferably with the subject refraining from alcohol for 24h-48h.

People with diabetes >40years old, or diabetes >15 years duration and age >30 years, or with established macrovascular or microvascular disease are considered at high risk for cardiac disease (Framingham Risk Score >20%) People with prediabetes and diabetes (who do not meet the criteria above) should be screened following the recommendations outlined in Table 2: Approach on Who and How to Screen for Dyslipidemia.

Dyslipidemia treatment recommendations are outlined in Table 3: Pharmacological Treatment Recommendations and Targets

The primary treatment goal for people with diabetes is LDL-C ≤2.0 mmol/L, which is generally achievable with statin monotherapy.

Urine Albumin-to-Creatinine Ratio (ACR) & Serum At diagnosis of type 2 diabetes or 5 years after diagnosis of Creatinine (eGFR) type 1 diabetes and yearly thereafter. As the ACR can be elevated with recent major exercise, fever, urinary tract infection, congestive heart failure, menstruation or acute severe elevations of blood pressure (BP) or blood glucose (BG), screening for albuminuria should be delayed in the presence of these conditions. Intravascular volume contraction e.g. dehydration or any acute illness can transiently lower kidney function, and GFR estimation for screening purposes should be delayed until such conditions resolve. If ACR >20.0 mg/mmol (macroalbuminuria) this is indicative of chronic kidney disease (CKD). RN or RD should refer to the PCP. If eGFR \leq 60 ml/min OR ACR \geq 2.0 mg/mmol (microalbuminuria) and there is no established diagnosis of CKD order serum creatinine for eGFR in 3 months AND 2 repeat random urine ACRs performed over the next 3 months. If eGFR < 60mL/min or 2 or 3 ACRs > 2.0 mg/mmol (indicative of chronic kidney disease) refer to PCP. If ACR and/or eGFR is indicative of CKD. It is recommended that a urine dipstick test be performed (by the PCP), either in the laboratory or at point of care, as a screen for renal disease other than diabetic nephropathy. People with diabetes and CKD should have a random urine ACR and a serum creatinine converted into an eGFR performed at least every 6 months Cobalamin (Vitamin B₁₂) At least every one to two years in patients on long-term treatment with Metformin¹. If vitamin B12 levels are below range, please discuss with PCP

Table 1: Approach on Who to Screen for Dyslipidemia

Who to screen

Men 40 years of age or older; women 40 years of age or older (or postmenopausal)

- Consider earlier in ethnic groups at increased risk such as South Asian or indigenous individuals All patients with any of the following conditions, regardless of age
- · Clinical evidence of atherosclerosis
- · Abdominal aortic aneurysm
- · Diabetes mellitus
- Arterial hypertension
- · Current cigarette smoking
- · Stigmata of dyslipidemia (corneal arcus, xanthelasma, xanthoma)
- · Family history of premature CVD*
- · Family history of dyslipidemia
- Chronic kidney disease (eGFR ≤ 60 mL/min/1.73 m² or ACR ≥ 3 mg/mmol)
- Obesity (BMI ≥ 30)
- · Inflammatory diseases (RA, SLE, PsA, AS, IBD)
- · HIV infection
- · Erectile dysfunction
- · COPD
- · History of hypertensive disorder of pregnancy

ACR, albumin-to-creatinine ratio; AS, ankylosing spondylitis; BMI, body mass index; COPD, chronic obstructive pulmonary disease; CVD, cardiovascular disease; eGFR, estimated glomerular filtration rate; IBD, inflammatory bowel disease; PsA, psoriatic arthritis; RA, rheumatoid arthritis; SLE, systemic lupus erythematous.

Men younger than 55 years of age and women younger than 65 years of age in first-degree relatives. Data from Anderson et al.¹

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Table 2: Approach on How to Screen for Dyslipidemia

How to screen

For all

- · History and physical examination
- Standard lipid profile*: TC, LDL-C, HDL-C, non-HDL-C,[†] TG
- FPG or A1c
- eGFR
- · Lipoprotein(a)—once in patient's lifetime, with initial screening

Optional

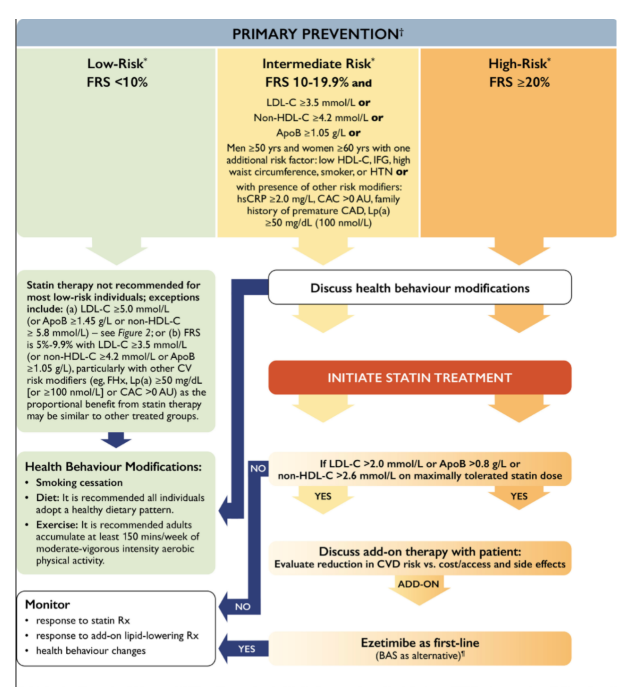
- ApoB
- Urine ACR (if eGFR <60 mL/min/1.73 m², hypertension, or diabetes)

A1c, glycated hemoglobin; ACR, albumin-to-creatinine ratio; ApoB, Apolipoprotein B; eGFR, estimated glomerular filtration rate; FPG, fasting plasma glucose; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; TC, total cholesterol; TG, triglycerides.

* Nonfasting lipid testing is recommended in most adults for screening; however, for individuals with a history of triglyceride levels > 4.5 mmol/L, measurement of fasting lipid levels are recommended. † It is now generally preferable to follow non-HDL-C or ApoB levels over LDL-C when interpreting lipid results, particularly when TG are ≥ 1.5 mmol/L.

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Table 3: Pharmacological Treatment Indications and Targets



PStatin indicated conditions consists of all documented ASCVD conditions, as well as other high-risk primary prevention conditions in the absence of ACSVD, such as most patients with diabetes, those with chronic kidney disease and those with a LDL-C ≥5.0 mmol/L.

Calculate risk using the Framingham Risk Score (FRS) — refer to the iCCS available on the App Store or on Google Play

"Screening should be repeated every 5 years for men and women aged 40 to 75 years using the modified FRS or CLEM to guide therapy to reduce major CV events. A risk assessment might also be completed whenever a patient's expected risk status changes.

 \P studies have evaluated the efficacy of BAS for the prevention of ASCVD, but results have been inconclusive.

FRS = Framingham risk score; LDL-C = low-density lipoprotein cholesterol; HDL-C = high-density lipoprotein cholesterol; ApoB = apolipoprotein B; IFG = impaired fasting glucose; HTN = hypertension hsCRP = high-sensitivity C-reactive protein; CAC = coronary artery calcium; AU – Agatston unit; Rx = prescription; BAS = bile acid sequestrant

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Table 4: Indications and Contraindications for Prescribing Diabetes Supplies

	Indications	Contraindications			
Prescribe Capillary Blood Glucometer & glucometer test strips	To assess glycemic control from blood glucose results in response to non-insulin anti-hyperglycemic agents, insulin and lifestyle management, quality control activities and patient teaching The results are used to determine if a patient is euglycemic, hyperglycemic or hypoglycemic so appropriate interventions and education can occur	 The patient or substitute decision maker refuses to monitor capillary blood glucose The patient is unable to monitor capillary blood glucose due to physical or cognitive limitations Considerations should be made for patients who are unable to monitor due to financial constraints SMBG not recommended due to DC guidelines, but is ultimately up to RN or RD clinical judgement 			
Prescribe Flash Glucometer & sensors	 To assess glycemic control from interstitial fluid glucose results in response to non-insulin antihyperglycemic agents, insulin and lifestyle management, quality control activities and patient teaching The results are used to determine if a patient is euglycemic, hyperglycemic or hypoglycemic so appropriate interventions and education can occur 	 The patient or substitute decision maker refuses to monitor flash interstitial glucose The patient is unable to monitor flash interstitial glucose due to physical or cognitive limitations Considerations should be made for patients who are unable to monitor due to financial constraints Flash Glucose monitoring not recommended due to DC guidelines, but is ultimately up to RN or RD clinical judgement The patient develops skin irritation or other reactions in response to the sensor 			
To assess blood ketone level patients with Type 1 Diabeted. The results are used to determine test strips To assess blood ketone patients with Type 1 Diabeted. The results are used to determine the patient is at risk for Dialeted. Ketoacidosis so appropriate interventions and education occur.		 The patient or substitute decision maker refuses to monitor blood ketone levels The patient is unable to monitor blood ketone levels due to physical or cognitive limitations Considerations should be made for patients who are unable to monitor due to financial constraints 			
Prescribe lancets & needle tips for insulin pens	Insulin pen needles or syringes for patients injecting insulin or GLP1ra	The length of the needles should be determined based on the current best practice recommendations for injections			

See Program Folders/Diabetes/Procedures/DEP - 12 Prescribing Diabetes Supplies and/or Making Medication Changes in Practice Solutions

Appendix D:

Sample Lab Requisition

Nan	Ontario Ministry of Health and Long-Term Car Laboratory Requisit Requisitioning Clinicine	tion	Lab	oratory Use Only				
Pa	uline Pariser							
Add	Iress							
	D Bay Street, Suite 300, PO Bo	ox 5,						
	ronto, ON,	- ·	Clin	ician/Practitioner's Contact Number for Urg	ent Resul	ts	Service Date	
	G 1N8	•	1	440 \ 000 4000			yyyy mm dd	
Clin	ician/Practitioner Number	CPSO / Registration No.	_``	416 / 960-1366 Ext.	Version	Sex	Date of Birth	
	9822	_				X M	yyyy mm dd	
	ock (✓) one:		Prov	vince Other Provincial Registration Number			Patient's Telephone Contact Number	
	OHIP/Insured	ninsured WSIB					416 204-1256	
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No	te: Separate requisitions are rec	uired for cytology, his	tolo	gy / pathology and tests performed	by Publ	ic He	ealth Laboratory	
×	Biochemistry		x	Hematology		X	Viral Hepatitis (check one only)	
	Glucose Random	∑ Fasting	100	CBC	, cate a	4:D5-10 42	Acute Hepatitis	
O	HbA1C	A doming	\vdash	Prothrombin Time (INR)		\dashv	Chronic Hepatitis	
XX	Creatinine (eGFR)			Immunology		\vdash	Immune Status / Previous Exposure	
	Uric Acid			Pregnancy Test (Urine)		\vdash	Specify: Hepatitis A	
	Sodium		Н	Mononucleosis Screen			Hepatitis B	
\vdash	Potassium		Т	Rubella			Hepatitis C or order individual hepatitis tests in the	
\vdash	Chloride		\vdash	Prenatal: ABO, RhD, Antibody Screen			"Other Tests" section below	
	СК		1	(titre and ident. if positive)		Pro	ostate Specific Antigen (PSA)	
\vdash	ALT			Repeat Prenatal Antibodies			otal PSA Free PSA	
	Alk. Phosphatase		Microbiology ID & Sensitivities (if warranted)			Specify one below: Insured – Meets OHIP eligibility criteria		
\vdash	Bilirubin							
	Albumin	113-7.	20 100 100 100 100 100 100 100 100 100 1		ninsured - Screening: Patient responsible for payment			
	Lipid Assessment (includes Choleste	erol, HDL-C, Triglycerides,		Vaginal		Vit	amin D (25-Hydroxy)	
X	calculated LDL-C & Chcl/HDL-C ration be ordered in the "Other Tests" section	on of this form)		Vaginal / Rectal – Group B Strep		□ In	nsured - Meets OHIP eligibility criteria:	
X	Albumin / Creatinine Ratio, Urine			Chlamydia (specify source):		_	osteopenia; osteoporosis; rickets; renal disease; malabsorption syndromes;	
	Urinalysis (Chemical)			GC (specify source):			medications affecting vitamin D metabolism	
	Neonatal Bilirubin:			Sputum		الا	Ininsured - Patient responsible for payment	
	Child's Age: days	hours		Throat		Ot	her Tests - one test per line	
	Clinician/Practitioner's tel. no.			Wound (specify source):			4-74-74-74-74-74-74-74-74-74-74-74-74-74	
	Patient's 24 hr telephone no.		\vdash	Urine				
	Therapeutic Drug Monitoring:		1	Stool Culture				
	Name of Drug #1	-	╀	Stool Ova & Parasites				
	Name of Drug #2		\vdash	Other Swabs / Pus (specify source):		_		
		hr. #2 hr.	0-	Silmon Collection (Authorities	. S. SHINE			
		hr. #2 hr.	Tim	ecimen Collection Date	HESCASTIC.			
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I hereby certify the tests ordered are not for registered in or out patients of a hospital.			cal Occult Blood Test (FOBT) (check FOBT (non CCC) ColonCar poratory Use Only		K FOB	T (CCC) no other test can be ordered on this form		
Dahart Swith DD CDE								
	obert Smith RD CDE	MD16						
As X	s per medical directive TCFHT-	-MD16 02/09/2014						
	nician/Practitioner Signature	Date						
			_				7620.450	

Appendix E

Implementer Competency Checklist for Prescribing Diabetes Supplies

Implementer Name:	
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CDA Guidelines Chapter Reviews						
Diabetes Canada 2018 Clinical Practice Guidelines for the Prevention and Management of Diabetes in Canada	Date Reviewed	Signature				
Diabetes and Driving						
Appendix 5: Self-Monitoring of Blood Glucose (SMBG) Recommendation Tool for Healthcare Providers						
Fit Forum for Injection Technique Canada. (2020). Fit Forum for Injection Technique Canada: Recommendations for Best Practice in Injection Technique.						
Canadian Insulin Injection Recommendations: FIT Technique Plus, 2020.						