



Medicines & Healthcare products Regulatory Agency

NovoRapid PumpCart in the Roche Accu-Chek Insight insulin pump: risk of insulin leakage causing hyperglycaemia and diabetic ketoacidosis

Date of Issue:	26-May-22	Ref	erence No:	NatPSA/2022/004/MHRA
This alert is for action	by: All Healthcare institutions	provid	ing specialist diabe	etes services to patients
	d complex National Patient Safety Al hisations without executive boards)			coordinated by an executive leader or diabetes and heads of procurement.
Explanation of iden	ified safety issue:	A	ctions required	
associated with insulin PumpCart prefilled insu Insight Insulin pump. In consequences arising fi including diabetic ketoa Although the manufactur has implemented a num to reduce the incidence these is inconclusive ar protect patients. The main reasons for th 1. The adapter ne cartridge at an bending of the leakage from th cases; 2. Cracked cartrid cracks, that ma such as droppir approximately 3 Risk minimisation strate improvements to the pu communications on how cartridge. An enhanced implemented in Septerr continue to be reported The manufacturer has r (FSN) on this issue, wh to this alert. Patients sh of following the advice i This includes using the tubing for certain lot nur To protect patient safety professionals should inf Chek Insight of the risk	arer Roche Diabetes Care (RDC) her of risk minimisation strategie of these events, the impact of ad we are taking further action to he leakages were: edle being inserted in the insulin incorrect angle, which may cause heedle and subsequent insulin he septum, in approximately 70% ges, including non-visible hairling y result from mechanical shocks ing the cartridge or pump; in 30% of cases; egies have included technical imp and adapter and safety v to handle the pump and prefille design of adaptor and tubing wa aber 2021 but events of leakage , albeit at reduced rates. eleased a <u>Field Safety Notice</u> ich should be followed in addition ould be informed of the importar n the manufacturer's latest <u>FSN</u> . new versions of adapter and mbers. y, diabetes healthcare form patients who use the Accu- of leakage. Clinical care decision ure patients are moved onto	s 2. of 3. of 4. s 5. ce A	Accu-Chek Ins in your organis provided them care. Any affe quarantined. Contact users undertake a pa assessment (s to determine s alternative pur Identify suitab use local proc acquire them. Onboard patie ensure an app per standard p initiating pump organisation. Patients with o use affected d suitable altern this has been the patient-cer continued use any stage of th alert requires a be completed additional info	ents to new pumps and propriate follow-up period as practice and guidance for therapy within your diabetes can continue to devices if there is a lack of ative insulin therapy or if deemed necessary after ntred consultation. Any of the affected device at he implementation of this a local risk assessment to and documented (see

For any enquiries about this alert contact: info@mhra.gov.uk

Additional information:

Information on adverse incidents

Cases with serious clinical consequences describing leakages of insulin, including cracked cartridges, in association with Accu-Chek Insight Pump and NovoRapid PumpCart Cartridges have been reported. In both 2020 and 2021, 25 serious cases each year (including cases where a patient required urgent medical treatment or hospitalisation) were reported to the MHRA in association with an insulin leakage event in UK patients, including 18 cases and 17 cases respectively of DKA.

So far in 2022 one case of DKA and 2 additional serious cases of hyperglycaemia have been reported to the MHRA in association with leakage events. In addition, non-serious cases of hyperglycaemia resulting from inadequate insulin supply have been also reported. In most cases, users did not require urgent medical intervention.

Risk assessment

A risk assessment must be recorded with all users. This should involve a discussion of the risks of continuing treatment with the affected device and consider the best interest of the patient and the management of their diabetes.

Patients presenting with unexplained hyperglycaemia identified as due to 'set failure' is a known risk experienced by users of insulin pumps. The issue associated with the Accu-Chek Insight pumps is different and is due to leakages directly from the reservoir, either by insulin escaping through cracks in the glass cartridge wall, or the cartridge septum not providing an adequate seal at the point of connection to the cannula. The leaked insulin can pool within the pump itself. As such, the problem may not be identified as quickly as other leaks since the user may not be aware of the leakage. If unidentified, the interrupted insulin delivery may lead to life-threatening consequences.

Effective training in the use of insulin pumps and continued vigilance of a patient's clinical presentation, combined with regular blood glucose monitoring, will reduce the likelihood of events such as hyperglycaemia and DKA. There is a risk that continued use of the Accu-Chek Insight pump with the prefilled glass cartridge may lead to leakages, particularly if the instructions for use are not followed closely.

The use of multiple daily injections (MDI) insulin regimens or other non-pump insulin delivery devices while patients await the onboarding of a new insulin pump should only be considered if suitable training and support is provided and where the benefits outweigh the risks associated with discontinuation of the Accu-Chek Insight pump.

If the risk assessment indicates the patient's condition is best managed by continuing therapy with the Accu-Chek Insight pump, instruct patients not to use cartridges that have been dropped, even if there are no visible cracks. Cracks may develop over time following a drop or other mechanical shock. Inform users of the importance of following the advice in the manufacturer's latest <u>FSN</u>. This includes using the new adapter and tubing and checking the pump and cartridge regularly.

All versions of the Accu-Chek Insight pumps are affected by this action. To note, RDC ceased marketing the devices in the UK at the end of 2021 and new patients will not be offered the pump. Existing patients remaining on the Accu-Chek Insight pump will be fully supported until the end of their warranty only. The risk assessment should include a plan to move to an alternative therapy delivery device before the end of the warranty period.

Link to useful resources

NICE guidelines on issuing pumps

Stakeholder engagement:

This action has been endorsed by the Commission on Human Medicines (CHM) and its Expert Advisory Groups and the Device Expert Advisory Committee (DEAC).

MHRA have consulted with NHS England and NHS Improvement, and representatives from the Scottish, Welsh, and Northern Ireland Governments. MHRA have also conducted patient engagement activities with users of insulin pumps.

Please check <u>website</u> for when actions should be ceased or advice to check for date restriction are lifted.







Llywodraeth Cymru Welsh Government

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