

April 1, 2024

Dear Health Care Providers:

RE: End of Humalog® Saskatchewan Biosimilars Initiative Transition Period

As you may be aware, the Saskatchewan Biosimilars Initiative was announced on October 20, 2022. Coverage for several reference biologics has already transitioned to biosimilar versions. The six-month transition period for Humalog® 100 units/mL started on October 1, 2023.

Starting April 1, 2024, Humalog® 100 units/mL is no longer eligible for coverage under the Saskatchewan Drug Plan.

- Patients will be responsible for the full cost of Humalog® 100 units/mL products, unless they have an approved exemption.
- Patients will need to use a biosimilar version of insulin lispro to maintain Saskatchewan Drug Plan coverage of their treatment.

The table below shows the available biosimilar insulin formats for insulin lispro:

| Insulin | Reference Biologic Brand and Formats | Biosimilar Insulin Brand and Formats | End of Transition Period |
|--------------------------------|---|---|--------------------------|
| Insulin lispro 100 units/mL | HUMALOG - Cartridge - Pre-filled pen - Vial | ADMELOG - Cartridge - Pre-filled pen - Vial | March 31, 2024 |

See the enclosed medSask insulin comparison chart for more details.

*Please note: Coverage of **Humalog® 200 units/mL** will continue to be available for patients who need a higher concentration formula, as there is no equivalent biosimilar format at this time.*

Please visit www.saskatchewan.ca/biosimilars for the most up-to-date information on the Saskatchewan Biosimilars Initiative.

Insulin Pump Users:

- The biosimilar (Admelog®) is compatible with various insulin pump models from Insulet (Omnipod), Medtronic, Tandem, and Ypsomed.
- Contact the insulin pump manufacturer for questions about insulin compatibility with specific insulin pump models.

One-Time Fill of Reference Biologic Medications:

- Starting April 1, 2024, patients who require an **urgent** refill of Humalog® 100 units/mL and have not yet transitioned to a covered biosimilar may be eligible for a *one-time fill* of the reference biologic medication.
- A one-time fill is intended to prevent treatment gaps or delays related to the patient's ability to transition to a biosimilar before their next refill.
- **Pharmacists should call** the Biosimilars Initiative team at **1-800-667-2549, option 3**, and leave a message to determine if their patient may be eligible for a one-time fill.
- If eligible for a one-time fill of the reference biologic, the patient will pay for the reference biologic according to their usual co-payment and/or deductible. Patients will have until their next refill to coordinate a biosimilar transition with their prescriber or pharmacist.
- **Reminder:** Pharmacists can support patients to transition to a biosimilar insulin without a prescription.

Exemptions:

- The Saskatchewan Biosimilars Initiative includes a provision for exemptions to the policy where patients must remain on the reference biologic for medical reasons.
- Please consider all contributing factors prior to requesting an exemption. The medSask tool to assess unexpected responses to biosimilars may be helpful (enclosed).
- The current Exemption Request Form is available on our webpage, www.saskatchewan.ca/biosimilars, under section 6, [Prescriber Forms](#).

Questions and Support:

For support with the Saskatchewan Biosimilars Initiative policy and drug coverage:

- Visit www.saskatchewan.ca/biosimilars
- Email sk.biosimilars@health.gov.sk.ca
- Call the Saskatchewan Drug Plan: 1-800-667-7581 (306-787-3317 in Regina)
 - To request a one-time fill, pharmacists should call 1-800-667-2549 (option 3).

For clinical support from medSask:

- **Health care providers:**
 - Visit medsask.usask.ca/professional-practice/biosimilars-in-Saskatchewan
 - Email druginfo@usask.ca
 - Call 1-800-667-3425 (306-966-6340 in Saskatoon)
- **Patients:**
 - Visit medsask.usask.ca/general-public/biosimilars-in-saskatchewan
 - Email med.sask@usask.ca
 - Call 1-800-665-3784 (306-966-6378 in Saskatoon)

INSULIN LISPRO

Comparison Chart for Individuals Switching to Biosimilar Insulin

Rapid acting insulin analogue, bolus/prandial and for use in subcutaneous pump system



- Use same dose (unit to unit) when transitioning to biosimilar.
- No clinical differences in onset, peak, or duration of action.
- No expected differences in adverse effects.

| Product | Humalog® (reference biologic) | | | Admelog® (biosimilar) | |
|--|---|-----------------------------------|---------------------------------|---|---------------------------------|
| Strength | 100 units/mL [#] | | | 100 units/mL | |
| Manufacturer | Eli Lilly | | | Sanofi-Aventis | |
| DIN | 02229705 | 02403412 | 02470152 | 02469898 | 02469871 |
| Supplied As | Cartridge—for use with HumaPen Savvio® or HumaPen Luxura® HD reusable pen (discontinued) | Prefilled pen—KwikPen® | Prefilled pen—Junior KwikPen® | Cartridge—for use with AllStar® Pro or JuniorSTAR® reusable pen | Prefilled pen—SoloSTAR® |
| Pen colour, injection button colour | HumaPen Savvio®: graphite or red pen HumaPen Luxura® HD: green pen | Dark blue pen, burgundy dose knob | Dark blue pen, blue dose knob | AllStar® Pro, JuniorSTAR®: blue or silver pen | Yellow pen, burgundy button |
| Administration | Remind individuals that the 'feel' of delivery devices may be different, but the basic mechanics of dialing a dose and subcutaneously injecting the insulin with a pen remain the same. | | | | |
| Dosing Increments | HumaPen Savvio®: 1-60 units in 1 unit increments HumaPen Luxura® HD: ½-30 units in ½ unit increments | 1-60 units in 1 unit increments | ½-30 units in ½ unit increments | AllStar® Pro: 1-80 units in 1 unit increments JuniorSTAR®: 1-30 units in ½ unit increments | 1-80 units in 1 unit increments |
| "Clicks" as dose delivered | ✓ | X | X | AllStar® Pro: ✓ JuniorSTAR®: ✓ | ✓ |
| End of dose "click" | X | X | X | X | X |
| Prevents dialing of more units than remain | HumaPen Savvio®: ✓ HumaPen Luxura® HD: X | ✓ | ✓ | AllStar® Pro: ✓ JuniorSTAR®: X | ✓ |
| Needle Compatibility | All pen devices are compatible with BD Pro Ultra-fine™, Insupen®, NovoFine® and Unifine® pen needles. | | | | |
| Storage: unopened | Refrigerated (2-8 °C) until expiration date. Keep away from direct heat and light. Do not freeze. | | | | |
| Storage: in use | Room temperature [^] (max. 30 °C) for up to 28 days. Keep away from direct heat and light. Do not freeze. | | | | |
| Ok to return to fridge when in use? | X | X | X | X | X |

Transitioning To A Biosimilar: Assessment Of Unexpected Response

- Biosimilars are demonstrated to be as effective and safe as the reference biologic. The unexpected response may not be related to the use of a biosimilar.
- When assessing an unexpected response: acknowledge and address patient concerns, use objective measures in addition to subjective information, and consider all potential factors.

| FACTORS | CONSIDERATIONS FOR REVIEW |
|--|--|
| <p>Drug Storage Deviations from manufacturer recommended storage may compromise efficacy.</p> | <ul style="list-style-type: none"> • Storage conditions <ul style="list-style-type: none"> ✓ Not exposed to temperature extremes (including during transport) ✓ Storage time at room temperature not exceeded ✓ Drug not expired |
| <p>Drug Regimen Non-adherence may lead to treatment failure resulting in unnecessary changes to or escalation of treatment.</p> | <ul style="list-style-type: none"> • Adherence <ul style="list-style-type: none"> ✓ Original reference biologic discontinued by patient ✓ Administered dose is the same as the reference biologic ✓ Dose given on time and as scheduled (i.e., no interruption of therapy) |
| <p>Drug Administration Improper use of the device could result in delivery of subtherapeutic dose.</p> | <ul style="list-style-type: none"> • Site of administration <ul style="list-style-type: none"> ✓ Appropriate and different from last site of administration • Dose delivery (as applicable) <ul style="list-style-type: none"> ✓ Plunger of prefilled syringe completely depressed ✓ Viewing window indicates complete drug delivery ✓ Autoinjector held in place at least 10 seconds ✓ Dose not accidentally discharged (i.e., autoinjector button pressed too soon) |
| <p>Drug Interactions Concomitant medications or supplements may:</p> <ul style="list-style-type: none"> • reduce efficacy of the biosimilar; • increase side effects; or • have side effects that mimic a disease flare. | <ul style="list-style-type: none"> • New use of: <ul style="list-style-type: none"> • Prescription medications • Over-the-counter medications • Supplements • Samples • Products ordered on the internet or purchased outside of Canada |
| <p>Clinical Status Of Condition Being Treated</p> | <ul style="list-style-type: none"> • Natural disease progression • Possibility of disease flare |

| FACTORS | CONSIDERATIONS FOR REVIEW |
|--|--|
| <p>Other Therapies Used To Manage Condition</p> | <ul style="list-style-type: none"> • Adherence or recent changes to: <ul style="list-style-type: none"> • Concomitant medications • Nonpharmacologic management (e.g. physical therapy, psychotherapy, diet, exercise, sleep/rest, etc.) |
| <p>Overall Health Status</p> | <ul style="list-style-type: none"> • Change in physical health including comorbid conditions, injury, or new diagnosis • Change in mental, emotional, social health (e.g. financial instability, work stress, access to care, etc.) |
| <p>Nocebo Effect</p> <p>Negative expectations may influence treatment outcomes.</p> | <ul style="list-style-type: none"> • Patient knowledge about biosimilars and sources of information • Patient anxiety about transitioning to the biosimilar • Health care provider confidence in the quality, safety, and efficacy of biosimilars |

Managing Injection Site Pain

The transition to a different product may result in a change to how the injection feels. Consider:

- Product formulation factors: excipients, pH, volume, temperature, viscosity
- Device features: needle length and gauge
- Injection technique: injection speed and movement during injection
- Patient factors: low body weight, female sex, mental health status, disease severity, expectations of pain

Unexpected and severe adverse effects should be reported to Health Canada:

<https://www.canada.ca/en/health-canada/services/drugs-health-products/medeffect-canada/adverse-reaction-reporting.html>



Who to contact with questions or concerns:

- Saskatchewan Biosimilars Initiative: email sk.biosimilars@health.gov.sk.ca or call 1.800.667.2549 (306.787.8744 in Regina), option 3.
- medSask: email druginfo@usask.ca or call 1.800.667.3425 (306.966.6340 in Saskatoon)