

URGENT VOLUNTARY RECALL: Pharmacy Level, June 18, 2025

Levothyroxine Sodium Tablets USP 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg and 175 mcg

Accord Healthcare, Inc. ("Accord Healthcare") is voluntarily recalling nine lots of Levothyroxine Sodium Tablets USP 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg and 175 mcg, at the Pharmacy Level.

This recall is being initiated because out of specification results were observed during long term stability testing of Levothyroxine Sodium Tablets USP for 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg and 175 mcg. The out of specification results were observed on an assay conducted on nine lots of Levothyroxine Sodium Tablets.

The assay content of Levothyroxine Sodium Tablets for lots D2300325 (25 mcg, 94.8% at 24 months), D2400536 (25 mcg, 94.4% at 12 months), D2400679 (25 mcg, 94.4% at 12 months), D2300087 (50 mcg, 94.9% at 24 months), D2300092 (100 mcg, 93.2% at 24 months), D2400722 (100 mcg, 94.7% at 12 months), D2300104 (112 mcg, 93.8% at 24 months), D2300076 (150 mcg, 94.7% at 24 months), and D2300042 (175 mcg, 94.9% at 24 months) was observed below the approved specification range of 95.0%—105.0% set forth on the label. This means that the observed level of active ingredient in the product was below the approved specification.

These may affect the efficacy of the medication. As patient safety is the highest priority, Accord Healthcare is taking immediate action to recall the affected product lots.

Please examine your inventory of Accord Healthcare's Levothyroxine Sodium Tablets USP 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg and 175 mcg for the below listed lot numbers carefully.

The product label for the recalled product should have the following details, please also refer to the enclosed product labels included with this recall letter.

Strength	Bottle Pack size	Product NDC	Lot No.	Expiry Date
25 mcg	90 Tablets	16729-447-15	D2300325	01/2026
25 mcg	90 Tablets	16.729-447-15	D2400536	02/2026
25 mcg	1000 Tablets	16729-447-17	D2400679	02/2026
50 mcg	90 Tablets	16729-448-15	D2300087	12/2025
100 mcg	1000 Tablets	16729-451-17	D2300092	12/2025
100 mcg	1000 Tablets	16729-451-17	D2400722	03/2026
112 mcg	1000 Tablets	16729-452-17	D2300104	12/2025
150 mcg	1000 Tablets	16729-455-17	D2300076	12/2025
175 mcg	1000 Tablets	16729-456-17	D2300042	12/2025





Pharmacy - Please perform the following activities:

- Examine your inventory immediately for the listed lot numbers of Levothyroxine Sodium Tablets USP 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg and 175 mcg.
- Immediately discontinue distribution of the recalled Lot numbers of Accord Healthcare's Levothyroxine Sodium Tablets USP 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg and 175 mcg.
- Promptly complete the attached Product Recall Response Form and reply even if you have <u>NO</u> Product to return.
- If you do have product to return, complete the attached Product Recall Response Form, quarantine the stock and follow the instructions given on the recall response form.
- If you have further distributed this lot number to other retailers, please immediately contact them and advise them of the recall and have them return their outstanding recalled stock to you. Return this stock as per the instructions on the attached Product Recall Response Form.

Your assistance is appreciated and necessary to prevent any potential health risk to the consumer.

Accord is working with Inmar Inc. to complete this recall. This recall is being made with the knowledge of the Food & Drug Administration. Your cooperation and prompt response to this notice is much appreciated.

<u>Please complete and return the enclosed "PRODUCT RECALL RESPONSE FORM" as soon as possible, but no later than five business days from receipt of this letter.</u>

Completed Product Recall Response form should be emailed, or sent via FAX to INMAR.

Attn: Inmar Rx Solutions, 3845 Grand Lakes Way, Grand Prairie, TX 75050.

INMAR Email: rxrecalls@inmar.com. FAX: 1-817-868-5362.

If you have any questions about the logistics for returning affected lot or other issues, please call Recall Services at 1-855-297-6127, Monday – Friday (excluding holidays), 9am to 5 pm EST.

INMAR will send you a <u>Return Goods Authorization</u> and <u>shipping label</u>. Appropriate credit for the returned product plus handling and shipping expenses will be issued to you upon receipt of the recalled product with the completed Return Goods Authorization. All recalled products returned without a Return Goods Authorization may delay the issuance of your credit.

We appreciate your assistance in this matter.

Sincerely,

Sabita Nair, RAC, ASQ-CPGP Vice President – Regulatory Affairs Accord Healthcare, Inc. 8041 Arco Corporate Drive, Suite 200

Raleigh, NC 27617, USA



Each tablet contains 25 mcg (0.025 mg) levothyroxine sodium USP.

See package insert for full prescribing information. Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture.

Dispense in a tight, light-resistant container as described in USP.

Do not accept if seal over bottle opening is broken or missing.

Manufactured for: Accord Healthcare, Inc., Raleigh, NC 27617.

Manufactured by: Intas Pharmaceuticals Limited, Camp Road, Selaqui, Dehradun-248 197, INDIA.

NDC 16729-447-15

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Each tablet contains 25 mcg (0.025 mg) levothyroxine sodium USP.

See package insert for full prescribing information.

Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture

Dispense in a tight, light-resistant container as described in USP. Do not accept if seal over bottle opening is broken or missing. Manufactured for: Accord Healthcare, Inc., Raleigh, NC 27617.

Manufactured by: Intas Pharmaceuticals Limited, Camp Road, Selagui, Dehradun-248 197, INDIA

NDC 16729-447-17

Levothyroxine

Sodium Tablets, USP

25 meg (0.025 mg)

Rx Only

1000 Tablets

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Each tablet contains 50 mcg (0.05 mg) levothyroxine sodium USP.

See package insert for full prescribing information. Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture.

Dispense in a tight, light-resistant container as described in USP.

Do not accept if seal over bottle opening is broken or missing.

Manufactured for: Accord Healthcare, Inc., Raleigh, NC 27617.

Manufactured by: Intas Pharmaceuticals Limited, Camp Road, Selaqui, Dehradun-248 197, INDIA NDC 16729-448-15

Levothyroxine Sodium Tablets, USP 50 mcg (0.05 mg) Rx Only 90 Tablets Accord

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Each tablet contains 100 meg (0.1 mg) levothyroidne sodium USP. See package insert for full prescribing information.

Contains FD&C Yellow No. 5 (tartrazine) as a color additive.

Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture.

Dispense in a tight, light-resistant container as described in USP. Do not accept if seal over bottle opening is broken or missing. Manufactured for: Accord Healthcare, Inc.,

Raleigh, NC 27617. Manufactured by: Intas Pharmaceuticals Limited. Camp Road, Setaqui, Dehradun-248 197, INDIA.

NDC 16729-451-17

Levothyroxine Sodium Tablets, USP

100 mca (0.1 mg)

Rx Only

1000 Tablets

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Each tablet contains 112 mcg (0.112 mg) levothyroxine sodium USP. See package insert for full prescribing information.

Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture.

Dispense in a tight, light-resistant container as described in USP. Do not accept if seal over bottle opening is broken or missing.

opening is broken or missing.
Manufactured for:
Accord Healthcare, Inc.,
Raleigh, NC 27617.
Manufactured by:
Intas Pharmaceuticals Limited,
Camp Road, Selaqui,

Dehradun-248 197, INDIA.

NDC 16729-452-17

Levothyroxine Sodium Tablets, USP

112 mgg (0.112 mg)

Rx Only

1000 Tablets

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Each tablet contains 150 mcg (0.15 mg) levothyroxine sodium USP.

See package insert for full prescribing information.

Store at 20°C to 25°C (68°F to \mathcal{T} 0°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture.

Dispense in a tight, light-resistant container as described in USP.
Do not accept if seal over bottle opening is broken or missing.
Manufactured for:
Accord Healthcare, inc.,

Raleigh, NC 27617. Manufactured by: Intas Pharmaceuticals Limited, Camp Road, Selaqui, Dehradun-248 197, INDIA. NDC 16729-455-17

Levothyroxine Sodium Tablets, USP

150 meg (0.15 mg)

Rx Only

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1000 Tablets



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Each tablet contains 175 mgg (0.175 mg) levothyroxine sodium USP. See package insert for full prescribing information.

Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) (see USP Controlled Room Temperature). Protect from light and moisture.

Dispense in a tight, light-resistant container as described in USP. Do not accept if seal over bottle opening is broken or missing.

Manufactured for: Accord Healthcare, Inc.. Raleigh, NC 27617. Manufactured by: Intas Pharmaceuticals Limited, Camp Road, Selaqui,

Dehradun-248 197, INDIA

NDC 16729-456-17

Levothyroxine Sodium Tablets, USP

175 meg (0.175 mg)

Rx Only

1000 Tablets

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PRODUCT RECALL RESPONSE FORM

Product Recall Date: June 18, 2025

Voluntary Recall: Pharmacy Level

Strength	Bottle Pack size	Product NDC	Lot No.	Expiry Date
25 mcg	90 Tablets	16729-447-15	D2300325	01/2026
25 mcg	90 Tablets	16729-447-15	D2400536	02/2026
25 mcg	1000 Tablets	16729-447-17	D2400679	02/2026
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100 mcg	1000 Tablets	16729-451-17	D2300092	12/2025
100 mcg	1000 Tablets	16729-451-17	D2400722	03/2026
112 mcg	1000 Tablets	16729-452-17	D2300104	12/2025
150 mcg	1000 Tablets	16729-455-17	D2300076	12/2025
175 mcg	1000 Tablets	16729-456-17	D2300042	12/2025

<u>Please fill out this form completely.</u> By doing so, this will acknowledge that you have read and understand the recall instructions and have taken the appropriate action.

Customer Name	DEA #	
	ided the processing of your form will be delayed.	
Address		
City	State	
Zip		
Contact Name (please print)	Telephone #	
Contact Signature	Date	
If you did not purchase the product directly	from the Manufacturer, please complete the follo	wing section.
Purchased from: Name	DEA #	
Address		
City	State	
Zip		





Please check all appropriate boxes: ☐ I have read and understand the recall instructions provided in the letter. ☐ I have checked my stock and have quarantined inventory consisting of bottles/units. Any adverse events associated with recalled product? □Yes □ NO If yes, please explain: Please describe your business: I have checked my stock and: Do not have any stock of recalled items. OR Have quarantined and listed in the box above the quantity of bottles/units of Levothyroxine Sodium Tablets USP 25 mcg, 50 mcg, 100 mcg, 112 mcg, 150 mcg, 175 mcg and will be returning them to Inmar, as soon as possible. Upon receipt of this Response Form, Inmar will issue return authorization label(s). Please indicate the number of box labels needed:___ Completed Product Recall Response form should be emailed, or sent via FAX to INMAR, Attn: Inmar Rx Solutions, 3845 Grand Lakes Way, Grand Prairie, TX 75050. INMAR Email: rxrecalls@inmar.com. FAX: 1-817-868-5362. Even if you do not possess any inventory of the lot being recalled, we would appreciate it if you could still fill out and return the "PRODUCT RECALL RESPONSE FORM". If you have any questions about the logistics for returning affected lot or other issues, please call Recall

Services at 1-855-297-6127, Monday - Friday (excluding holidays), 9am to 5 pm EST.

