

Subcutaneous Amivantamab and Hyaluronidase-Ipuj: Dosing and Administration

SC Amivantamab Indications and Recommended Dosing Schedule¹

Adults with locally advanced or metastatic NSCLC with <i>EGFR</i> exon 19 deletions or exon 21 L858R substitution mutations		Adults with locally advanced or metastatic NSCLC with <i>EGFR</i> exon 20 insertion mutations	
1L amivantamab + lazertinib	2L amivantamab + carboplatin and pemetrexed	1L amivantamab + carboplatin and pemetrexed	2L amivantamab single agent
Q2W/Q4W	Q3W	Q3W	Q2W/Q4W

SC Amivantamab is Supplied at a Concentration of 160 mg/mL with 2,000 units Hyaluronidase/mL in Single-Dose Vials as Follows:¹



10 mL vial (1,600 mg amivantamab + 20,000 units hyaluronidase)



14 mL vial (2,240 mg amivantamab + 28,000 units hyaluronidase)

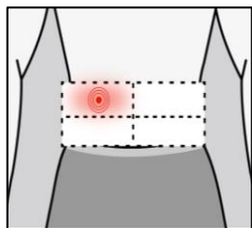


15 mL vial (2,400 mg amivantamab + 30,000 units hyaluronidase)

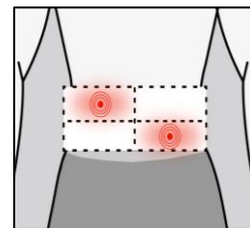


22 mL vial (3,520 mg amivantamab + 44,000 units hyaluronidase)

SC Amivantamab Important Administration Information¹



- Administer SC amivantamab in the abdomen over ≈5 minutes to minimize injection site irritation
- Do not inject into tattoos or scars or areas where the skin is red, bruised, tender, hard, not intact or within 2 inches (5 cm) around the periumbilical area
- Rotate injection sites at the next scheduled dose

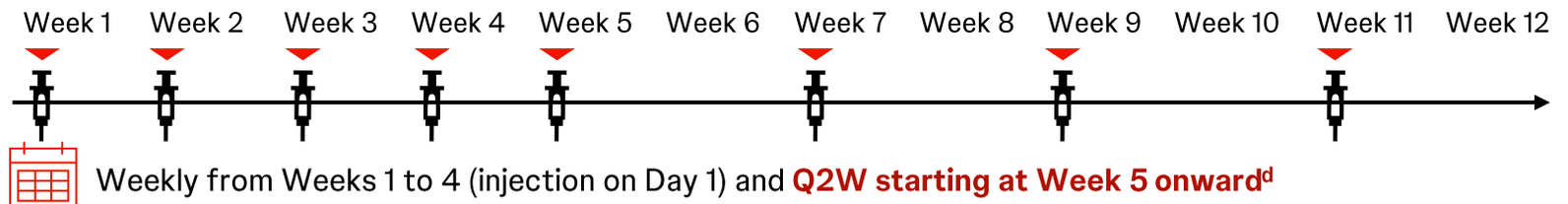


- If the total dose requires multiple injections of SC amivantamab, administer each injection consecutively in separate quadrants of the abdomen; do not exceed 15 mL in each syringe
- If the patient experiences pain, pause or slow delivery; if the pain is not alleviated by pausing or slowing the delivery rate, deliver the rest of the dose in a second injection site on the opposite side of the abdomen

SC Amivantamab Dosing in Combination With Lazertinib or as a Single Agent¹

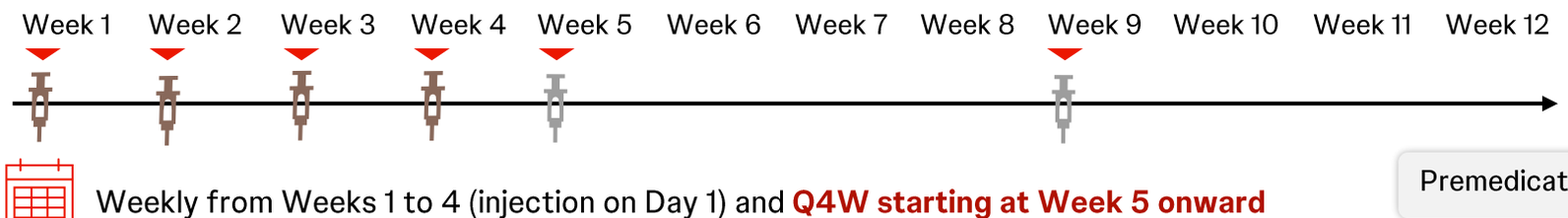
SC amivantamab Q2W^a

	<80 kg		≥80 kg	
All weeks	1,600 mg	10 mL required	2,240 mg	14 mL required



SC amivantamab Q4W^a

	<80 kg		≥80 kg	
Weeks 1-4	1,600 mg	10 mL required	2,240 mg	14 mL required
Week 5+	3,520 mg	22 mL ^b required	4,640 mg	29 mL ^{b,c} required



Premedications →

Administer SC amivantamab any time after lazertinib when given on the same day. See the Prescribing Information for lazertinib for recommended lazertinib dosing information and drug interactions. See the Prescribing Information for SC amivantamab for dosage and administration, including prophylactic anticoagulation, premedication, and prophylactic/concomitant dermatologic medication information.

^aDose adjustments not required for subsequent body weight changes. ^bDivide the dose volume approximately equally into 2 syringes (each syringe should not exceed 15 mL). ^cFor the 29 mL dose volume, use one 14 mL vial and one 15 mL vial to minimize waste. If a different combination of vials is used, discard unused portion. ^dMay switch to SC amivantamab Q4W at next scheduled dose on or after Week 5. 1L, first-line; 2L, second-line; EGFR, epidermal growth factor receptor; NSCLC, non-small cell lung cancer; Q2W, every 2 weeks; Q3W, every 3 weeks; Q4W, every 4 weeks; SC, subcutaneous.



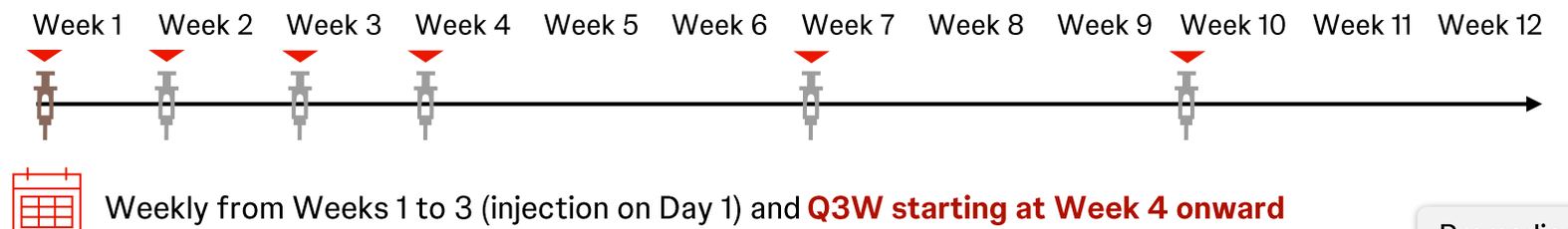
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Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

SC Amivantamab Dosing in Combination With Carboplatin and Pemetrexed¹

SC amivantamab Q3W^a

<80 kg			≥80 kg		
Week 1	1,600 mg	10 mL required	Week 1	2,240 mg	14 mL required
Week 2+	2,400 mg	15 mL required	Week 2+	3,360 mg	21 mL ^b required



Premedications →

Please refer to the Prescribing Information for SC amivantamab for order of administration and regimen for amivantamab in combination with carboplatin and pemetrexed. Refer to the Prescribing Information of pemetrexed and carboplatin for complete information regarding appropriate use, including Boxed Warnings for carboplatin. See the Prescribing Information for SC amivantamab for dosage and administration, including premedication and prophylactic/concomitant dermatologic medication information.

Contraindications¹

SC amivantamab is contraindicated in patients with known hypersensitivity to hyaluronidase or to any of its excipients. Please refer to the Prescribing Information for SC amivantamab for additional safety information.

Amivantamab IV to SC Transition¹⁻⁷

Considerations for Amivantamab IV to SC transition:

SC Prescribing Information:

The company cannot support any practices, procedures, or dosage administration techniques that deviate from the approved product labeling. Please refer to the DOSAGE AND ADMINISTRATION, DOSAGE FORMS AND STRENGTHS, CONTRAINDICATIONS (hypersensitivity to hyaluronidase), WARNINGS AND PRECAUTIONS, ADVERSE REACTIONS, USE IN SPECIFIC POPULATIONS, and CLINICAL STUDIES sections of the Prescribing Information for SC amivantamab for complete information.¹

Ongoing clinical trial:

Based on PK noninferiority from PK simulations comparing IV to SC systemic exposure from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2, patients treated with IV amivantamab may transition to SC amivantamab administration.²⁻⁴ PALOMA-2 (NCT05498428) is an ongoing, phase 2, open-label, international, parallel cohort study evaluating the efficacy and safety of SC amivantamab with chemotherapy and/or lazertinib in patients with *EGFR*-mutated locally advanced or metastatic NSCLC.⁵⁻⁷



^aDose adjustments not required for subsequent body weight changes. ^bDivide the dose volume approximately equally into 2 syringes (each syringe should not exceed 15 mL). EGFR, epidermal growth factor receptor; IV, intravenous; NSCLC, non-small cell lung cancer; PK, pharmacokinetic; Q3W, every 3 weeks; SC, subcutaneous.

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Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

Amivantamab IV to SC Transition^{1,2}

The tables below outline the recommended IV and SC starting dosages which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2}

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹
Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for Q2W IV amivantamab and Q2W SC amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	700 mg	No dose	1,050 mg	No dose
Weeks 2-4	1,050 mg	1,600 mg	1,400 mg	2,240 mg
Week 5 (may switch per PI)	1,050 mg	1,600 mg	1,400 mg	2,240 mg
Week 6	No dose			
Week 7 and Q2W thereafter	1,050 mg	1,600 mg	1,400 mg	2,240 mg

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q4W at their next scheduled dose on or after Week 5.¹
Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for Q2W IV amivantamab and Q4W SC amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV (Q2W)	SC (Q4W)	IV (Q2W)	SC (Q4W)
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	700 mg	No dose	1,050 mg	No dose
Weeks 2-4	1,050 mg	1,600 mg	1,400 mg	2,240 mg
Week 5 (may switch per PI)	1,050 mg	3,520 mg	1,400 mg	4,640 mg
Week 5 and all doses onwards	1,050 mg (Q2W)	3,520 mg (Q4W)	1,400 mg (Q2W)	4,640 mg (Q4W)

Adult patients currently receiving IV amivantamab Q3W may switch to SC amivantamab Q3W at their next scheduled dose on or after Week 4.¹
Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for Q3W IV amivantamab and Q3W SC amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	1,050 mg	No dose	1,400 mg	No dose
Weeks 2-3	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Week 4 (may switch per PI)	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Weeks 5 and 6	No dose			
Week 7 and Q3W thereafter	1,750 mg	2,400 mg	2,100 mg	3,360 mg

Click Links Below for Information on Recommended Dose Reductions, Dose Modifications, and Management for ARs with Amivantamab^{1,2}

[IV amivantamab dose reductions](#) →

[SC amivantamab dose reductions](#) →

[Hypersensitivity & ARRs](#) →

[ILD/pneumonitis](#) →

[VTEs](#) →

[Dermatologic ARs](#) →

[Other ARs](#) →



AR, adverse reaction; ARR, administration-related reaction; ILD, interstitial lung disease; IV, intravenous; PI, prescribing information; Q2W, every 2 weeks; Q3W, every 3 weeks; Q4W, every 4 weeks; SC, subcutaneous; VTE, venous thromboembolic events.

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Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

SC Amivantamab Indications and Recommended Dosing Schedule¹

Adults with locally advanced or metastatic NSCLC with *EGFR* exon 19 deletions or exon 21 L858R substitution mutations

Adults with locally advanced or metastatic NSCLC with *EGFR* exon 20 insertion mutations

1L amivantamab + lazertinib

2L amivantamab + carboplatin and pemetrexed

1L amivantamab + carboplatin and pemetrexed

2L amivantamab single agent

Q2W/Q4W

Q3W

Q3W

Q2W/Q4W

Premedications

SC Q2W/Q4W dosing schedule →

SC Q3W dosing schedule →

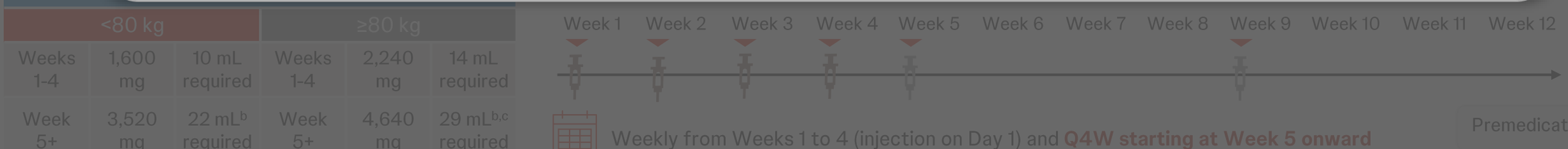


Prior to the initial injection of SC amivantamab, administer premedications as described below. Glucocorticoid administration is required at the initial dose at Week 1 Day 1 only, and upon re-initiation after prolonged dose interruptions, then as necessary for subsequent injections. Administer both antihistamine and antipyretic prior to all SC amivantamab doses.

Medication	Dose	Route of administration	Dosing window prior to SC amivantamab administration
Antihistamine ^a	Diphenhydramine (25 mg to 50 mg) or equivalent	Intravenous	15 to 30 minutes
		Oral	30 to 60 minutes
Antipyretic ^a	Acetaminophen (650 mg to 1,000 mg) or equivalent	Intravenous	15 to 30 minutes
		Oral	30 to 60 minutes
Glucocorticoid ^b	Dexamethasone (20 mg) or equivalent	Intravenous	45 to 60 minutes
		Oral	At least 60 minutes
Glucocorticoid ^c	Dexamethasone (10 mg) or equivalent	Intravenous	45 to 60 minutes
		Oral	60 to 90 minutes

^aRequired at all doses. ^bRequired at initial dose (Week 1, Day 1) or at the next subsequent dose in the event of an administration-related reaction. ^cOptional for subsequent doses.

1. RYBREVANT FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026.



Administer SC amivantamab any time after lazertinib when given on the same day. See the Prescribing Information for lazertinib for recommended lazertinib dosing information and drug interactions. See the Prescribing Information for SC amivantamab for dosage and administration, including prophylactic anticoagulation, premedication, and prophylactic/concomitant dermatologic medication information.

^aDose adjustments not required for subsequent body weight changes. ^bDivide the dose volume approximately equally into 2 syringes (each syringe should not exceed 15 mL). ^cFor the 29 mL dose volume, use one 14 mL vial and one 15 mL vial to minimize waste. If a different combination of vials is used, discard unused portion. ^dMay switch to SC amivantamab Q4W at next scheduled dose on or after Week 5. 1L, first-line; 2L, second-line; EGFR, epidermal growth factor receptor; NSCLC, non-small cell lung cancer; Q2W, every 2 weeks; Q3W, every 3 weeks; Q4W, every 4 weeks; SC, subcutaneous.

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Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

Amivantamab IV to SC Transition^{1,2}

The tables below outline the recommended IV and SC starting dosages which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2}

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹
Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for Q2W IV amivantamab and Q2W SC amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC

IV Amivantamab Dose Reductions^{1,a}



Initial dose	1,050 mg	1,400 mg	1,750 mg	2,100 mg
1st dose reduction	700 mg	1,050 mg	1,400 mg	1,750 mg
2nd dose reduction	350 mg	700 mg	1,050 mg	1,400 mg
3rd dose reduction	⚠ Discontinue			

^aRecommended dosage modifications for ARs for IV amivantamab in combination with lazertinib: when administering IV amivantamab in combination with lazertinib, if there is an AR requiring dose reduction after withholding treatment and resolution, reduce the dose of IV amivantamab first. See the lazertinib Prescribing Information for information about dosage modifications for lazertinib. Recommended dosage modifications for ARs for IV amivantamab in combination with carboplatin and pemetrexed: when administering IV amivantamab in combination with carboplatin and pemetrexed, modify the dosage of ≥1 drug. Withhold or discontinue IV amivantamab as shown in Table 7 of the Prescribing Information. Refer to the Prescribing Information for carboplatin and pemetrexed for additional dosage modification information.

AR, adverse reaction; IV, intravenous.

1. RYBREVANT® (amivantamab-vmjw) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026.

Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	1,050 mg	No dose	1,400 mg	No dose
Weeks 2-3	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Week 4 (may switch per PI)	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Weeks 5 and 6			No dose	
Week 7 and Q3W thereafter	1,750 mg	2,400 mg	2,100 mg	3,360 mg

Click Links Below for Information on Recommended Dose Reductions, Dose Modifications, and Management for ARs with Amivantamab^{1,2}

- [IV amivantamab dose reductions](#)
- [SC amivantamab dose reductions](#)
- [Hypersensitivity & ARRs](#)
- [ILD/pneumonitis](#)
- [VTEs](#)
- [Dermatologic ARs](#)
- [Other ARs](#)



AR, adverse reaction; ARR, administration-related reaction; ILD, interstitial lung disease; IV, intravenous; PI, prescribing information; Q2W, every 2 weeks; Q3W, every 3 weeks; Q4W, every 4 weeks; SC, subcutaneous; VTE, venous thromboembolic events.

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Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

Amivantamab IV to SC Transition^{1,2}

The tables below outline the recommended IV and SC starting dosages which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2}

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹
Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for
Q2W IV amivantamab and
Q2W SC amivantamab^{1,2}

<80 kg

≥80 kg

IV

SC

IV

SC

SC Amivantamab Dose Reductions^{1,a}

X

Initial dose	1,600 mg ^b	2,240 mg ^c	2,400 mg ^d	3,360 mg ^e	3,520 mg ^f	4,640 mg ^g
1st dose reduction	1,050 mg ^h	1,600 mg ⁱ	1,600 mg ⁱ	2,240 mg ^j	2,400 mg ^k	3,360 mg ^l
2nd dose reduction	700 mg ^m	1,050 mg ^h	1,050 mg ^h	1,600 mg ⁱ	1,600 mg ⁱ	2,240 mg ^j
3rd dose reduction	⚠ Discontinue					

^aRecommended dosage modifications for ARs for SC amivantamab in combination with lazertinib: when administering SC amivantamab in combination with lazertinib, if there is an AR requiring dose reduction after withholding treatment and resolution, reduce the dose of SC amivantamab first. See the lazertinib Prescribing Information for information about dosage modifications for lazertinib. Recommended dosage modifications for ARs for SC amivantamab in combination with carboplatin and pemetrexed: when administering SC amivantamab in combination with carboplatin and pemetrexed, modify the dosage of ≥1 drug. Withhold or discontinue SC amivantamab as shown in Table 7 of the Prescribing Information. Refer to the Prescribing Information for carboplatin and pemetrexed for additional dosage modification information. ^bDose volume 10 mL. ^cDose volume 14 mL. ^dDose volume 15 mL. ^eDose volume 21 mL. ^fDose volume 22 mL. ^gDose volume 29 mL. ^hDose volume 6.6 mL. ⁱDose volume 10 mL. ^jDose volume 14 mL. ^kDose volume 15 mL. ^lDose volume 21 mL. ^mDose volume 4.4 mL. AR, adverse reaction; SC, subcutaneous.

1. RYBREVANT FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026.

Weeks 2-3	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Week 4 (may switch per PI)	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Weeks 5 and 6			No dose	
Week 7 and Q3W thereafter	1,750 mg	2,400 mg	2,100 mg	3,360 mg

Click Links Below for Information on Recommended Dose Reductions, Dose Modifications, and Management for ARs with Amivantamab^{1,2}

[IV amivantamab dose reductions](#) →

[SC amivantamab dose reductions](#) →

[Hypersensitivity & ARRs](#) →

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[VTEs](#) →

[Dermatologic ARs](#) →

[Other ARs](#) →

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AR, adverse reaction; ARR, administration-related reaction; ILD, interstitial lung disease; IV, intravenous; PI, prescribing information; Q2W, every 2 weeks; Q3W, every 3 weeks; Q4W, every 4 weeks; SC, subcutaneous; VTE, venous thromboembolic events.

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1. RYBREVANT FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026. 2. RYBREVANT® (amivantamab-vmjw) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2025.

Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

Amivantamab IV to SC Transition^{1,2}

The tables below outline the recommended IV and SC starting dosages which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2}

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹
Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for
Q2W IV amivantamab and
Q2W SC amivantamab^{1,2}

Hypersensitivity and ARRs¹



Recommended Dosage Modifications and Management for Hypersensitivity and ARRs for SC Amivantamab

Grade 1-2



INTERRUPT
SC amivantamab
injection if ARR
is suspected and
monitor patient
until reaction
symptoms
resolve



RESUME
injection upon
resolution of
symptoms



PREMEDICATE
Include corticosteroid
with premedications
for subsequent dose
(see Table 6 in
Prescribing
Information)

Grade 3



INTERRUPT
SC
amivantamab
injection



ADMINISTER
supportive
care
medications



MONITOR
patient
continuously
until reaction
symptoms
resolve



RESUME
injection upon
resolution of
symptoms



PREMEDICATE
Include
corticosteroid
with premedications
for subsequent dose
(see Table 6 in
Prescribing
Information)



**PERMANENTLY
DISCONTINUE**
SC
amivantamab
for recurrent
Grade 3 ARRs

Grade 4



**PERMANENTLY
DISCONTINUE**
SC amivantamab

Please see Warnings and Precautions (section 5.1) for further details regarding hypersensitivity and ARRs.

ARR, administration-related reaction; SC, subcutaneous.

1. RYBREVA FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026.

Recommended dosage for
Q3W IV amivantamab and
Q3W SC amivantamab^{1,2}

	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	1,050 mg	No dose	1,400 mg	No dose
Weeks 2-3	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Week 4 (may switch per PI)	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Weeks 5 and 6			No dose	
Week 7 and Q3W thereafter	1,750 mg	2,400 mg	2,100 mg	3,360 mg

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1. RYBREVA FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026. 2. RYBREVA® (amivantamab-vmjw) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2025.

Subcutaneous Amivantamab and Hyaluronidase-lpuj: Dosing and Administration

Amivantamab IV to SC Transition^{1,2}

The tables below outline the recommended IV and SC starting dosages which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2}

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹
 Amivantamab maximum trough concentration is typically observed at the end of the weekly dosing (Cycle 2 Day 1).¹

Recommended dosage for Q2W IV amivantamab and Q2W SC amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC

ILD/Pneumonitis¹



Recommended Dosage Modifications and Management for ILD/Pneumonitis for SC Amivantamab

Any grade



WITHHOLD
 SC amivantamab
 if ILD/pneumonitis
 is suspected



PERMANENTLY DISCONTINUE
 SC amivantamab
 if ILD/pneumonitis
 is confirmed

Please see Warnings and Precautions (section 5.2) for further details regarding ILD/pneumonitis.

ILD, interstitial lung disease; SC, subcutaneous.

1. RYBREVANT FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026.

Q3W IV amivantamab and Q3W SC amivantamab ^{1,2}	IV		SC	
	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	1,050 mg	No dose	1,400 mg	No dose
Weeks 2-3	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Week 4 (may switch per PI)	1,400 mg	2,400 mg	1,750 mg	3,360 mg
Weeks 5 and 6			No dose	
Week 7 and Q3W thereafter	1,750 mg	2,400 mg	2,100 mg	3,360 mg

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AR, adverse reaction; ARR, administration-related reaction; ILD, interstitial lung disease; IV, intravenous; PI, prescribing information; Q2W, every 2 weeks; Q3W, every 3 weeks; Q4W, every 4 weeks; SC, subcutaneous; VTE, venous thromboembolic events.

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1. RYBREVANT FASPRO™ (amivantamab and hyaluronidase-lpuj) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026. 2. RYBREVANT® (amivantamab-vmjw) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2025.

Subcutaneous Amivantamab Dosing and Administration

Amivantamab IV to SC Transition¹⁻⁵

The tables below outline the recommended IV and SC starting dosages, as well as the corresponding dose reductions, which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2} Based on PK noninferiority from PK simulations comparing IV to SC systemic exposures from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2 (NCT05498428), patients treated with IV amivantamab may transition to SC administration³⁻⁵

Recommended dosage for Q2W Amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	700 mg	No dose	1,050 mg	No dose

Venous Thromboembolic (VTE) Events¹



Recommended Dosage Modifications and Management for Venous Thromboembolic (VTE) Events for SC Amivantamab in Combination With Lazertinib

Grade 2-3



WITHHOLD SC amivantamab and lazertinib



ADMINISTER anticoagulation treatment as clinically indicated



Once anticoagulant treatment has been initiated, **RESUME SC amivantamab and lazertinib** at the same dose level, at the discretion of the treating physician

Grade 4 or recurrent Grade 2-3 despite therapeutic-level anticoagulation



WITHHOLD lazertinib and **PERMANENTLY DISCONTINUE** SC amivantamab



ADMINISTER anticoagulation treatment as clinically indicated



Once anticoagulant treatment has been initiated, **RESUME lazertinib** at the same dose level, at the discretion of the treating physician

Please see Warnings and Precautions (section 5.3) for further details regarding VTE events; applies to the combination with lazertinib.

SC, subcutaneous.

1. RYBREVANT FASPRO™ (amivantamab and hyaluronidase-lpui) [Prescribing Information]. Horsham, PA: Janssen Biotech, Inc.; 2026.

Considerations for Amivantamab IV to SC transition:¹⁻⁸



HCP clinical judgement:

HCP to use clinical judgement, patient's medical history & history with amivantamab, including prior hypersensitivity.



Important guidance on transition:

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹

Adult patients currently receiving IV amivantamab Q3W may switch to SC amivantamab Q3W at their next scheduled dose on or after Week 4.¹



SC Prescribing Information:

The company cannot support any practices, procedures, or dosage administration techniques that deviate from the approved product labeling. Please refer to the DOSAGE AND ADMINISTRATION, DOSAGE FORMS AND STRENGTHS, CONTRAINDICATIONS (hypersensitivity to hyaluronidase), WARNINGS AND PRECAUTIONS, ADVERSE REACTIONS, USE IN SPECIFIC POPULATIONS, and CLINICAL STUDIES sections of the Prescribing Information for SC Amivantamab for complete information.¹



Ongoing clinical trial:

Based on PK noninferiority from PK simulations comparing IV to SC systemic exposure from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2, patients treated with IV amivantamab may transition to SC amivantamab administration.³⁻⁵ PALOMA-2 (NCT05498428) is an ongoing, phase 2, open-label, international, parallel cohort study evaluating the efficacy and safety of SC amivantamab with chemotherapy and/or lazertinib in patients with EGFR-mutated locally advanced or metastatic NSCLC.⁶⁻⁸

¹Recommended dosage modifications for ARs for SC amivantamab in combination with lazertinib: when administering SC amivantamab in combination with lazertinib, if there is an AR requiring dose reduction after withholding treatment and resolution, reduce the dose of SC amivantamab first. See the lazertinib prescribing information for information about dosage modifications for lazertinib. Recommended dosage modifications for ARs for SC amivantamab in combination with carboplatin and pemetrexed: when administering SC amivantamab in combination with carboplatin and pemetrexed, modify the dosage of one or more drugs. Withhold or discontinue SC amivantamab as shown in Table 7 of the prescribing information. Refer to prescribing information for carboplatin and pemetrexed for additional dosage modification information. ²Starting dose volume is 10 mL. ³Starting dose volume is 14 mL. ⁴Starting dose volume is 15 mL. ⁵Starting dose volume is 21 mL. ⁶The dose volume should be 6.6 mL. ⁷The dose volume should be 4.4 mL. ⁸The dose volume should be 10 mL. ⁹The dose volume should be 14 mL. AR, adverse reaction; ARR, administration-related reaction; EGFR, epidermal growth factor receptor; IV, intravenous; NSCLC, non-small cell lung cancer; PK, pharmacokinetic; Q2W, every 2 weeks; Q3W, every 3 weeks; SC, subcutaneous; VTE, venous thromboembolism.

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Subcutaneous Amivantamab: Dosing and Administration

Amivantamab IV to SC Transition¹⁻⁵

The tables below outline the recommended IV and SC starting dosages, as well as the corresponding dose reductions, which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2} Based on PK noninferiority from PK simulations comparing IV to SC systemic exposures from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2 (NCT05498428), patients treated with IV amivantamab may transition to SC administration³⁻⁵

Recommended dosage for Q2W Amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	700 mg	No dose	1,050 mg	No dose

Dermatologic ARs¹



Recommended Dosage Modifications and Management for Dermatologic ARs (Including Dermatitis Acneiform, Pruritus, and Dry Skin) for SC Amivantamab

Grade 1-2



INITIATE
supportive care management as clinically indicated



REASSESS after 2 weeks; if rash does not improve, consider dose reduction

Grade 3



WITHHOLD
SC amivantamab and **INITIATE**
supportive care management as clinically indicated



Upon recovery to Grade ≤2, **RESUME**
SC amivantamab at reduced dose



If no improvement within 2 weeks, **PERMANENTLY DISCONTINUE**
treatment

Grade 4 or severe bullous, blistering, or exfoliating skin conditions^a



PERMANENTLY DISCONTINUE
SC amivantamab

Please see Warnings and Precautions (section 5.4) for further details regarding dermatologic ARs (including dermatitis acneiform, pruritus, dry skin, and TEN).

^aIncluding TEN.

AR, adverse reaction; SC, subcutaneous; TEN, toxic epidermal necrolysis.

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Considerations for Amivantamab IV to SC transition:¹⁻⁸



HCP clinical judgement:

HCP to use clinical judgement, patient's medical history & history with amivantamab, including prior hypersensitivity.



Important guidance on transition:

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹

Adult patients currently receiving IV amivantamab Q3W may switch to SC amivantamab Q3W at their next scheduled dose on or after Week 4.¹



SC Prescribing Information:

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Ongoing clinical trial:

Based on PK noninferiority from PK simulations comparing IV to SC systemic exposure from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2, patients treated with IV amivantamab may transition to SC amivantamab administration.³⁻⁵ PALOMA-2 (NCT05498428) is an ongoing, phase 2, open-label, international, parallel cohort study evaluating the efficacy and safety of SC amivantamab with chemotherapy and/or lazertinib in patients with EGFR-mutated locally advanced or metastatic NSCLC.⁶⁻⁸

^aRecommended dosage modifications for ARs for SC amivantamab in combination with lazertinib: when administering SC amivantamab in combination with lazertinib, if there is an AR requiring dose reduction after withholding treatment and resolution, reduce the dose of SC amivantamab first. See the lazertinib prescribing information for information about dosage modifications for lazertinib. Recommended dosage modifications for ARs for SC amivantamab in combination with carboplatin and pemetrexed: when administering SC amivantamab in combination with carboplatin and pemetrexed, modify the dosage of one or more drugs. Withhold or discontinue SC amivantamab as shown in Table 7 of the prescribing information. Refer to prescribing information for carboplatin and pemetrexed for additional dosage modification information. ^bStarting dose volume is 10 mL. ^cStarting dose volume is 14 mL. ^dStarting dose volume is 15 mL. ^eStarting dose volume is 21 mL. ^fThe dose volume should be 6.6 mL. ^gThe dose volume should be 4.4 mL. ^hThe dose volume should be 10 mL. ⁱThe dose volume should be 14 mL. AR, adverse reaction; ARR, administration-related reaction; EGFR, epidermal growth factor receptor; IV, intravenous; NSCLC, non-small cell lung cancer; PK, pharmacokinetic; Q2W, every 2 weeks; Q3W, every 3 weeks; SC, subcutaneous; VTE, venous thromboembolism.

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Subcutaneous Amivantamab: Dosing and Administration

Amivantamab IV to SC Transition¹⁻⁵

The tables below outline the recommended IV and SC starting dosages, as well as the corresponding dose reductions, which are provided in the current Prescribing Information for SC amivantamab and IV amivantamab.^{1,2} Based on PK noninferiority from PK simulations comparing IV to SC systemic exposures from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2 (NCT05498428), patients treated with IV amivantamab may transition to SC administration³⁻⁵

Recommended dosage for Q2W Amivantamab ^{1,2}	<80 kg		≥80 kg	
	IV	SC	IV	SC
Week 1 Day 1	350 mg	1,600 mg	350 mg	2,240 mg
Week 1 Day 2	700 mg	No dose	1,050 mg	No dose

Other ARs¹



Recommended Dosage Modifications and Management for Other ARs for SC Amivantamab

Grade 3



WITHHOLD
SC amivantamab
until recovery
to Grade ≤1
or baseline



RESUME
at the same dose if
recovery occurs
within 1 week



RESUME
at reduced dose if
recovery occurs
after 1 week but
within 4 weeks



**PERMANENTLY
DISCONTINUE**
if recovery does
not occur within
4 weeks

Grade 4



WITHHOLD
SC amivantamab
until recovery
to Grade ≤1
or baseline



RESUME
at reduced dose if
recovery occurs
within 4 weeks



**PERMANENTLY
DISCONTINUE**
if recovery does
not occur within
4 weeks



**PERMANENTLY
DISCONTINUE**
for recurrent
Grade 4 reactions

Please see Adverse Reactions (section 6.1) for further details regarding other ARs.

AR, adverse reaction; SC, subcutaneous.

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Considerations for Amivantamab IV to SC transition:¹⁻⁸



HCP clinical judgement:

HCP to use clinical judgement, patient's medical history & history with amivantamab, including prior hypersensitivity.



Important guidance on transition:

Adult patients currently receiving IV amivantamab Q2W may switch to SC amivantamab Q2W at their next scheduled dose on or after Week 5.¹

Adult patients currently receiving IV amivantamab Q3W may switch to SC amivantamab Q3W at their next scheduled dose on or after Week 4.¹



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Ongoing clinical trial:

Based on PK noninferiority from PK simulations comparing IV to SC systemic exposure from the phase 3 PALOMA-3 (NCT05388669) study, and IV to SC transition evaluated in PALOMA-2, patients treated with IV amivantamab may transition to SC amivantamab administration.³⁻⁵ PALOMA-2 (NCT05498428) is an ongoing, phase 2, open-label, international, parallel cohort study evaluating the efficacy and safety of SC amivantamab with chemotherapy and/or lazertinib in patients with EGFR-mutated locally advanced or metastatic NSCLC.⁶⁻⁸

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