Resource Guide

MVASI® is a biosimilar to Avastin® (bevacizumab), backed by Amgen expertise¹



INDICATIONS

MVASI® is a vascular endothelial growth factor inhibitor indicated for the treatment of:

MVASI[®], in combination with intravenous fluorouracil-based chemotherapy, is indicated for the first- or second-line treatment of patients with metastatic colorectal cancer (mCRC).

MVASI[®], in combination with fluoropyrimidine-irinotecan- or fluoropyrimidine-oxaliplatin-based chemotherapy, is indicated for the second-line treatment of patients with mCRC who have progressed on a first-line bevacizumab product-containing regimen.

Limitations of Use: MVASI® is not indicated for adjuvant treatment of colon cancer.

MVASI[®], in combination with carboplatin and paclitaxel, is indicated for the first-line treatment of patients with unresectable, locally advanced, recurrent or metastatic non-squamous non-small cell lung cancer (NSCLC).

MVASI® is indicated for the treatment of recurrent glioblastoma (GBM) in adults.

MVASI[®], in combination with interferon-alfa, is indicated for the treatment of metastatic renal cell carcinoma (mRCC).

MVASI[®], in combination with paclitaxel and cisplatin or paclitaxel and topotecan, is indicated for the treatment of patients with persistent, recurrent, or metastatic cervical cancer (CC).

Please see full Important Safety Information and click here for full Prescribing Information.

This brochure does not take the place of the reconstitution and preparation instructions located in the full Prescribing Information (PI). Please refer to the PI for specific instructions on preparing MVASI[®].



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Amgen can provide conversion support for institutions, including staff training, reimbursement assistance, and to facilitate an easier transition to MVASI[®].



IMPORTANT SAFETY INFORMATION

Serious adverse reactions (Warnings and Precautions)

- Serious and sometimes fatal adverse reactions with increased incidence in the bevacizumabtreated arm vs chemotherapy arm included:
 - Gastrointestinal (GI) perforation ranged from 0.3% to 3% of patients across clinical studies
 - Non-GI fistulae (<1% to 1.8%, highest in patients with cervical cancer)
 - Arterial thromboembolic events (Grade ≥3, 5%, highest in patients with GBM)
 - The incidence of wound healing and surgical complications, including serious and fatal complications, is increased in MVASI®-treated patients



Please see <u>full Important Safety Information</u> and click here for full Prescribing Information.

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SUPPLY AND STORAGE

SUPPLY¹

MVASI® is supplied as a sterile, colorless to pale yellow, preservative-free solution containing 25 mg/mL of bevacizumab-awwb in a single-dose vial. The vial stopper contains dry natural rubber.

Each carton of MVASI® contains either:

- 100 mg of MVASI® in 4 mL (25 mg/mL) (NDC 55513-206-01)
- 400 mg of MVASI® in 16 mL (25 mg/mL) (NDC 55513-207-01)

STORAGE¹

- Store at 2° to 8°C (36° to 46°F) in the original carton until time of use. MVASI® vials should be protected from light.
 - Diluted MVASI[®] solutions may be stored at 2° to 8°C (36° to 46°F) for up to 8 hours.



DO NOT FREEZE OR SHAKE.

Discard any unused portion remaining in the vial.

IMPORTANT SAFETY INFORMATION (cont'd)

Serious adverse reactions (Warnings and Precautions)

- Serious and sometimes fatal adverse reactions with increased incidence in the bevacizumab-treated arm vs chemotherapy arm included:
 - Hemorrhage (Grade 3–5) ranged from 0.4% to 7% of patients across clinical studies
 - Renal injury and proteinuria
 - Grade 3-4 proteinuria ranged from 0.7% to 7% in clinical studies
 - Nephrotic syndrome (<1%)

Please see <u>full Important Safety Information</u> and <u>click here for full Prescribing Information</u>.



DILUTION¹

Use appropriate aseptic technique. Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration.



• MVASI® is a colorless to pale yellow solution. Do not use vial if solution is cloudy, discolored, or contains particulate matter.



 Withdraw necessary amount of MVASI[®] and dilute in a total volume of 100 mL of 0.9% Sodium Chloride Injection, USP. Discard any unused portion left in a vial, as the product contains no preservatives.



• Do not administer or mix with dextrose solution.

ADMINISTRATION¹

Administer only as an intravenous (IV) infusion. Do not administer as an intravenous push or bolus.

- Do not initiate MVASI[®] until at least 28 days following major surgery. Administer MVASI[®] after the surgical incision has fully healed.
- First infusion: Administer over 90 minutes.
- Subsequent infusions: Administer over 60 minutes if first infusion is tolerated; administer all subsequent infusions over 30 minutes if infusion over 60 minutes is tolerated.

IMPORTANT SAFETY INFORMATION (cont'd)

- Additional serious adverse reactions with increased incidence in the bevacizumab-treated arm vs chemotherapy arm included:
 - Venous thromboembolism (Grade ≥3, 11% seen in GOG-0240)
 - Hypertension (Grade 3-4, 5%-18%)
 - Posterior reversible encephalopathy syndrome (PRES) (<0.5%)



Please see <u>full Important Safety Information</u> and click here for full Prescribing Information.

RECOMMENDED DOSES AND SCHEDULES

METASTATIC COLORECTAL CANCER (mCRC) ¹			
WITH IFL CHEMOTHERAPY	Every 2 Weeks 5 mg/kg		
WITH FOLFOX4	Every 2 Weeks 10 mg/kg		
WITH BEVACIZUMAB-AWWB + FLUOROPYRIMIDINE AND IRINOTECAN OR OXALIPLATIN	Every 2 Weeks 5 mg/kg OR Every 3 Weeks 7.5 mg/kg		
NON-SQUAMOUS NON-SMALL	CELL LUNG CANCER (NSCLC) ¹		
WITH CARBOPLATIN + PACLITAXEL	Every 3 Weeks 15 mg/kg		
RECURRENT GLIOI	BLASTOMA (rGBM) ¹		
AS SINGLE AGENT	Every 2 Weeks 10 mg/kg		
METASTATIC RENAL CE	LL CARCINOMA (mRCC) ¹		
WITH INTERFERON ALFA	Every 2 Weeks 10 mg/kg		
CERVICAL CANCER (CC) ¹			
WITH PACLITAXEL + CISPLATIN OR PACLITAXEL + TOPOTECAN	Every 3 Weeks 15 mg/kg		

IFL = irinotecan, leucovorin (folinic acid), and fluorouracil; FOLFOX4 = fluorouracil, leucovorin, and oxaliplatin.

• Patients should continue treatment until disease progression or unacceptable toxicity.¹

IMPORTANT SAFETY INFORMATION (cont'd)

- Additional serious adverse reactions with increased incidence in the bevacizumab-treated arm vs chemotherapy arm included:
 - Congestive heart failure (CHF): Grade ≥3 left ventricular dysfunction (1%)
 - Infusion-related reactions with the first dose of bevacizumab occurred in <3% of patients, and severe reactions occurred in 0.4% of patients

Please see <u>full Important Safety Information</u> and <u>click here for full Prescribing Information.</u>



CODING

NATIONAL DRUG CODES (NDCs)^{2,3}

BILLING		Each single-dose carton contains one vial of MVASI® (100 mg of bevacizumab- awwb) in 4 mL (25 mg/mL): NDC 55513-206-01
		Each single-dose carton contains one vial of MVASI® (400 mg of bevacizumab- awwb) in 16 mL (25 mg/mL): NDC 55513-207-01

METASTATIC COLORECTAL CANCER (mCRC)

	Malignant neoplasm of the following:		
ICD-10-CM⁴	CecumC18.0-C18.1Colon (various sites)C18.2-C18.9Rectosigmoid junctionC19Rectum, rectal ampullaC20Overlapping sites of rectum, anus, and anal canalC21.8		
HCPCS⁵	Q5107 injection, bevacizumab-awwb, 10 mg		
	96413: Chemotherapy administration, intravenous infusion technique; up to 1 hour, single or initial substance/drug.		
CPT ^{®6}	96415: Chemotherapy administration, intravenous infusion technique; each additional hour. Must be listed separately in addition to code for primary procedure.		
	96417: Chemotherapy administration, intravenous infusion technique; each additional sequential infusion (different substance/drug), up to one hour. Must be listed separately in addition to code for primary procedure.		

ICD = international classification of diseases; HCPCS = healthcare common procedure coding system; CPT = current procedural terminology.

IMPORTANT SAFETY INFORMATION (cont'd)

- Avoid use in patients with ovarian cancer who have evidence of recto-sigmoid involvement by pelvic examination or bowel involvement on CT scan or clinical symptoms of bowel obstruction
- Inform females of reproductive potential of the risk of ovarian failure prior to initiating treatment with MVASI[®]

Pregnancy warning

- Based on the mechanism of action and animal studies, MVASI® may cause fetal harm
- Advise female patients that MVASI® may cause fetal harm, and to inform their healthcare provider of a known or suspected pregnancy



Please see <u>full Important Safety Information</u> and click here for full Prescribing Information

6 click here for full Prescribing Information.

NON-SQUAMOUS NON-SMALL CELL LUNG CANCER (NSCLC)

	Malignant neoplasm of the following:		
	Trachea	C33	
	Bronchus and lung, main bronchus	C34.00-C34.02	
ICD-10-CM ⁴	Upper lobe, bronchus or lung	C34.10-C34.12	
	Middle lobe, bronchus or lung	C34.2	
	Lower lobe, bronchus or lung	C34.30-C34.32	
	Overlapping sites, bronchus or lung	C34.80-C34.82	
	Unspecified part, bronchus or lung	C34.90-C34.92	
HCPCS⁵	Q5107 injection, bevacizumab-awwb, 10 mg		
	96413: Chemotherapy administration, intravenous infusion technique; up to 1 hour, single or initial substance/drug.		
CPT ^{®6}	96415: Chemotherapy administration, intravenous infusion technique; each additional hour. Must be listed separately in addition to code for primary procedure.		
	96417: Chemotherapy administration, intravenous infusion technique; each additional sequential infusion (different substance/drug), up to one hour. Must be listed separately in addition to code for primary procedure.		

GLIOBLASTOMA

ICD-10-CM ⁴	Malignant neoplasm of the brain C71.0-71.9	
HCPCS⁵	Q5107 injection, bevacizumab-awwb, 10 mg	
	96413: Chemotherapy administration, intravenous infusion technique; up to 1 hour, single or initial substance/drug.	
CPT ^{®6}	96415: Chemotherapy administration, intravenous infusion technique; each additional hour. Must be listed separately in addition to code for primary procedure.	
	96417: Chemotherapy administration, intravenous infusion technique; each additional sequential infusion (different substance/drug), up to one hour. Must be listed separately in addition to code for primary procedure.	

IMPORTANT SAFETY INFORMATION (cont'd)

- Advise females of reproductive potential to use effective contraception during treatment with MVASI® and for 6 months after the last dose of MVASI®
- Advise nursing women not to breastfeed during treatment with MVASI® and for 6 months following their last dose of treatment
- MVASI[®] may impair fertility

Please see <u>full Important Safety Information</u> and <u>click here for full Prescribing Information.</u>



CODING (cont'd)

METASTATIC RENAL CELL CARCINOMA (mRCC)

	Malignant neoplasm of the following:		
ICD-10-CM⁴	Unspecified kidney, except renal pelvis C64	4.1-C64.2 4.9 5.1-C65.2, C65.9	
HCPCS⁵	Q5107 injection, bevacizumab-awwb, 10 mg		
	96413: Chemotherapy administration, intravenous infusion technique; up to 1 hour, single or initial substance/drug.		
96415: Chemotherapy administration, intravenous infusion technique; e additional hour. Must be listed separately in addition to code for primar			
	96417: Chemotherapy administration, intravenous infusion technique; each additional sequential infusion (different substance/drug), up to one hour. Must be listed separately in addition to code for primary procedure.		

PERSISTENT, RECURRENT, OR METASTATIC CARCINOMA OF THE CERVIX

	Malignant neoplasm of the following:		
ICD-10-CM⁴	Endocervix and exocervixC53.0-C53.1Overlapping sites of cervix uteri and unspecified sitesC53.8-C53.9of the cervix uteriC53.8-C53.9		
HCPCS⁵	Q5107 injection, bevacizumab-awwb, 10 mg		
	96413: Chemotherapy administration, intravenous infusion technique; up to 1 hour, single or initial substance/drug.		
96415: Chemotherapy administration, intravenous infusion technique; each additional hour. Must be listed separately in addition to code for primary prod			
	96417: Chemotherapy administration, intravenous infusion technique; each additional sequential infusion (different substance/drug), up to one hour. Must be listed separately in addition to code for primary procedure.		

IMPORTANT SAFETY INFORMATION (cont'd) Most common adverse reactions

- Across studies, the most common adverse reactions observed in bevacizumab patients at a rate >10% were:
 - Epistaxis
- Proteinuria
- Headache

• Rhinitis

- Hypertension
- Taste alteration
- Hemorrhage
- Lacrimation disorder
- Back pain
- Exfoliative dermatitis
- Across all studies, bevacizumab was discontinued in 8% to 22% of patients because of adverse reactions

• Dry skin

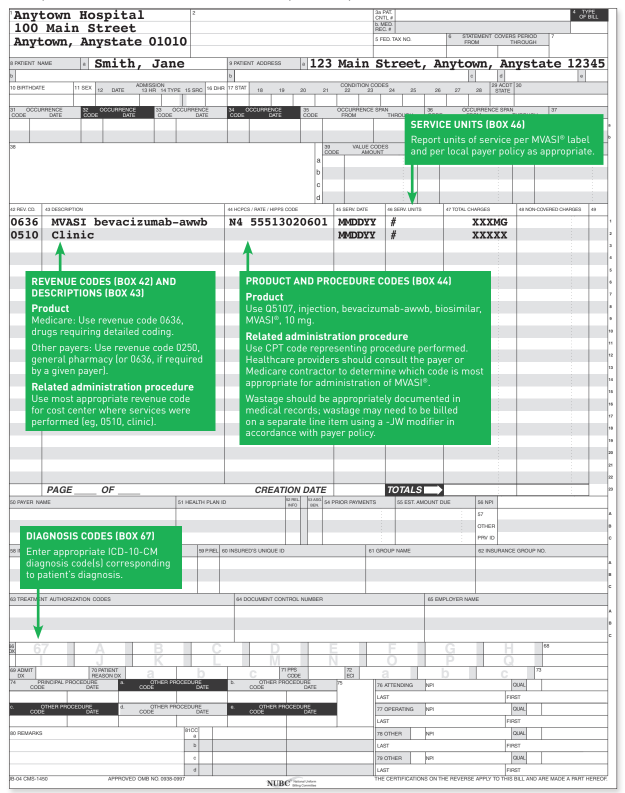


Please see **full Important Safety Information** and click here for full Prescribing Information.

HOSPITAL CODING FORM

The CMS 1450 for Hospital Outpatient

Sample UB-04 (CMS 1450) Form — Hospital Outpatient Administration



This sample form is intended as a reference for coding and billing for product and associated services. It is not intended to be directive; the use of the recommended codes does not guarantee reimbursement. Healthcare providers may deem other codes or policies more appropriate and should select the coding options that most accurately reflect their internal system guidelines, payer requirements, practice patterns, and the services rendered. Healthcare providers are responsible for ensuring the accuracy and validity of all billing and claims for appropriate reimbursement.



PHYSICIAN CODING FORM

The CMS 1500 for Physician Office

Sample CMS 1500 Form — Physician Office Administration

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	Л		CARRIER
	C) 02/12	PICA [
		ER 1a. INSURED'S I.D. NUMBER (For Program in Item 1)	
	Member ID#) (ID#) (ID#) (ID#)		
2. PATIENT'S NAME (Last Name, First Name, Middle Initial) Doe, John D	3. PATIENT'S BIRTH DATE SEX MM DD YY XX XX XX M F	4. INSURED'S NAME (Last Name, First Name, Middle Initial) Doe , John D	
5. PATIENT'S ADDRESS (No., Street)	6. PATIENT RELATIONSHIP TO INSURED	7. INSURED'S ADDRESS (No., Street)	
5555 Any Street	Self Spouse Child Other]	
CITY Anytown	AS 8. RESERVED FOR NUCC USE	CITY STATE	PATIENT AND INSURED INFORMATION
ZIP CODE TELEPHONE (Include Area Coc		ZIP CODE TELEPHONE (Include Area Code)	AMA
9. OTHER INSURED'S NAME (Last Name, First Name, Middle Initi		11. INSURED'S POLICY GROUP OR FECA NUMBER	B
9. OTHER INSORED'S NAME (Last Name, First Name, Middle Initi	a) 10.15 PATIENT'S CONDITION RELATED TO:	TLINSURED'S POLICY GROUP OR FECA NUMBER	
a. OTHER INSURED'S POLICY OR GROUP NUMBER	a. EMPLOYMENT? (Current or Previous)	a. INSURED'S DATE OF BIRTH SEX	URE
b. RESERVED FOR NUCC USE	b. AUTO ACCIDENT?		SN
	b. AUTO ACCIDENT? PLACE (Stat	b. OTHER CLAIM ID (Designated by NUCC)	AND
c. RESERVED FOR NUCC USE	c. OTHER ACCIDENT?	i c. INSURANCE PLAN NAME OR PROGRAM NAME	NT /
	YES NO		ATTE
d. INSURANCE PLAN NAME OR PROGRAM NAME	10d. CLAIM CODES (Designated by NUCC)	d. IS THERE ANOTHER HEALTH BENEFIT PLAN?	A -
PRODUCT CODE (BOX 24D)	ING & SIGNING THIS FORM.	13. INSURED'S OR AUTHORIZED PERSON'S SIGNATURE I authorize	
Use Q5107, injection, bevacizumab-awwl biosimilar, MVASI®, 10 mg.	b, he release of any medical or other information necessary ner to myself or to the party who accepts assignment	payment of medical benefits to the undersigned physician or supplier for services described below.	or
	DIAGN	NOSIS CODE (BOX 21)	
SIGNED	DATE DOCUN	nent appropriate ICD-10-CM	
14. DATE OF CURRENT ILLNESS, INJURY, or PREGNANCY (LM MM DD YY XX XX XX QUAL	QUAL: i i i anagina	bosis code(s) corresponding to	
17. NAME OF REFERRING PROVIDER OR OTHER SOURCE	1/a. diagno	t's diagnosis. Line A — primary SERVICES psis code. PP _ YY	
19. ADDITIONAL CLAIM INFORMATION (Designated by NUCC)	17b. NPI	NOSIS CODE (BOX 24E)	
		y diagnosis, from Box 21, relating to	
21. DIAGNOSIS OR NATURE OF ILLNESS OR INJURY Relate A-		CPT/HCPCS code listed in Box 24D.	
A. L XXX . XX B. L		23. PRIOR AUTHORIZATION NUMBER	
	G H K L	-	
	PROCEDURES, SERVICES, OR SUPPLIES (Explain Unusual Circumstances) DIAG NOS	F. G. H. I. J. DAYS EPSOT ID. RENDERING R \$CHARGES UNITS Partly DUAL PROVIDER ID. #	NOL
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1 N4 55513020601 XXXMG xx xx xx xx xx xx 11	05107 A	SERVICE UNITS (B	
			uld be appropriately
- xx xx xx xx xx xx 11	96XXX A	xxx xx # documented in med	dical records;
3	1 I I I I I I	wastage may need	to be billed on a using a -JW modifier
A		in accordance with	
	BOX 24D)		
5 Use CPT code represe	enting procedure	NPI	PHYSICIAN
performed. Healthca	re providers should Medicare contractor to		IS A
determine which code	a is most appropriate for	NPI	
25. FEDERAL TAX I.D. NUME administration of MVA		? 28. TOTAL CHARGE 29. AMOUNT PAID 30. Rsvd for NUCd \$ \$ \$ 1	C Use
31. SIGNATURE OF PHYSICIAN OR SUPPLIER 32. SER INCLUDING DEGREES OR CREDENTIALS (I certify that the statements on the reverse apply to this bill and are made a part thereof.)	VICE FACILITY LOCATION INFORMATION	33. BILLING PROVIDER INFO & PH # ()	
a.	NDI b.	a. NDI b.	
SIGNED DATE			2 12)
NUCC Instruction Manual available at: www.nucc.or	rg PLEASE PRINT OR TYPE	APPROVED OMB-0938-1197 FORM 1500 (0.	<u> </u>

This sample form is intended as a reference for coding and billing for product and associated services. It is not intended to be directive; the use of the recommended codes does not guarantee reimbursement. Healthcare providers may deem other codes or policies more appropriate and should select the coding options that most accurately reflect their internal system guidelines, payer requirements, practice patterns, and the services rendered. Healthcare providers are responsible for ensuring the accuracy and validity of all billing and claims for appropriate reimbursement.



MVASI® PRODUCT FACT SHEET

INDICATIONS

MVASI[®] is a vascular endothelial growth factor inhibitor indicated for the treatment of:

MVASI®, in combination with intravenous fluorouracil-based chemotherapy, is indicated for the first- or second-line treatment of patients with metastatic colorectal cancer (mCRC).

MVASI®, in combination with fluoropyrimidine-irinotecan- or fluoropyrimidine-oxaliplatin-based chemotherapy, is indicated for the second-line treatment of patients with mCRC who have progressed on a first-line bevacizumab product-containing regimen.

Limitations of Use: MVASI® is not indicated for adjuvant treatment of colon cancer.

MVASI[®], in combination with carboplatin and paclitaxel, is indicated for the first-line treatment of patients with unresectable, locally advanced, recurrent or metastatic non-squamous non-small cell lung cancer (NSCLC).

MVASI® is indicated for the treatment of recurrent glioblastoma (GBM) in adults.

MVASI®, in combination with interferon-alfa, is indicated for the treatment of metastatic renal cell carcinoma (mRCC).

MVASI®, in combination with paclitaxel and cisplatin or paclitaxel and topotecan, is indicated for the treatment of patients with persistent, recurrent, or metastatic cervical cancer (CC).

PRODUCT INFORMATION

NDC	Description	Quantity
55513-206-01	100 mg of MVASI® in 4 mL (25 mg/mL)	One per carton
55513-207-01	400 mg of MVASI® in 16 mL (25 mg/mL)	One per carton

STORAGE AND HANDLING REQUIREMENTS

Store at 2° to 8°C (36° to 46°F) in the original carton until time of use. MVASI[®] vials should be protected from light. DO NOT FREEZE OR SHAKE. Discard any unused portion remaining in the vial. Diluted MVASI[®] solutions may be stored at 2° to 8°C (36° to 46°F) for up to 8 hours. Store undiluted vials at 2° to 8°C (36° to 46°F) in the original carton until time of use. MVASI[®] vials should be protected from light.

SHIPPING CONTAINER INFORMATION

MVASI[®] should be unpacked and refrigerated.

MVASI® should not be stored in the shipping container.

PRODUCT EXPIRATION

The expiration date is printed on each dispensing pack and vial label.

SUPPLIED AND MARKETED BY

Amgen USA Inc.

amgen.com

MVASI.com

PRODUCT RETURNS

For information and instructions regarding product returns, please contact your wholesaler or Amgen Trade Operations at 1-800-28-AMGEN (1-800-282-6436). Credit for returns is subject to Amgen's current Product Return Policy.

PRODUCT INFORMATION

Medical Information: 1-800-77-AMGEN (1-800-772-6436)

REIMBURSEMENT INFORMATION

Amgen Assist 360™: 1-888-4ASSIST (1-888-427-7478) or www.AmgenAssistOnline.com



Please see **full Important Safety Information** and **click here for full Prescribing Information**.

IMPORTANT SAFETY INFORMATION

Serious adverse reactions (Warnings and Precautions)

- Serious and sometimes fatal adverse reactions with increased incidence in the bevacizumab-treated arm vs chemotherapy arm included:
 - Gastrointestinal (GI) perforation ranged from 0.3% to 3% of patients across clinical studies
 - Non-GI fistulae (<1% to 1.8%, highest in patients with cervical cancer)
 - Arterial thromboembolic events (Grade ≥3, 5%, highest in patients with GBM)
 - The incidence of wound healing and surgical complications, including serious and fatal complications, is increased in MVASI®-treated patients
 - Hemorrhage (Grade 3-5) ranged from 0.4% to 7% of patients across clinical studies
 - Renal injury and proteinuria
 - Grade 3–4 proteinuria ranged from 0.7% to 7% in clinical studies
 - Nephrotic syndrome (<1%)
- Additional serious adverse reactions with increased incidence in the bevacizumab-treated arm vs chemotherapy arm included:
 - Venous thromboembolism (Grade ≥3, 11% seen in GOG-0240)
 - Hypertension (Grade 3-4, 5%-18%)
 - Posterior reversible encephalopathy syndrome (PRES) (<0.5%)
 - Congestive heart failure (CHF): Grade ≥3 left ventricular dysfunction (1%)
 - Infusion-related reactions with the first dose of bevacizumab occurred in <3% of patients, and severe reactions occurred in 0.4% of patients
- Avoid use in patients with ovarian cancer who have evidence of recto-sigmoid involvement by pelvic examination or bowel involvement on CT scan or clinical symptoms of bowel obstruction
- Inform females of reproductive potential of the risk of ovarian failure prior to initiating treatment with MVASI®

Pregnancy warning

- Based on the mechanism of action and animal studies, MVASI® may cause fetal harm
- Advise female patients that MVASI® may cause fetal harm, and to inform their healthcare provider of a known or suspected pregnancy
- Advise females of reproductive potential to use effective contraception during treatment with MVASI® and for 6 months after the last dose of MVASI®
- Advise nursing women not to breastfeed during treatment with MVASI® and for 6 months following their last dose of treatment
- MVASI® may impair fertility

Most common adverse reactions

- Across studies, the most common adverse reactions observed in bevacizumab patients at a rate >10% were:
 - Epistaxis
 - Headache
 - Hypertension
 - Rhinitis
 - Proteinuria
 - Taste alteration
 - Dry skin
 - Hemorrhage
 - Lacrimation disorder
 - Back pain
 - Exfoliative dermatitis
- Across all studies, bevacizumab was discontinued in 8% to 22% of patients because of adverse reactions

Indication-specific adverse reactions

- In CC, Grade 3 or 4 adverse reactions in Study GOG-0240, occurring at a higher incidence (≥2%) in 218 patients receiving bevacizumab plus chemotherapy compared to 222 patients receiving chemotherapy alone, were abdominal pain (12% vs 10%), diarrhea (6% vs 3%), anal fistula (4% vs 0%), proctalgia (3% vs 0%), urinary tract infection (8% vs 6%), cellulitis (3% vs 0.5%), fatigue (14% vs 10%), hypertension (11% vs 0.5%), thrombosis (8% vs 3%), hypokalemia (7% vs 4%), hyponatremia (4% vs 1%), dehydration (4% vs 0.5%), neutropenia (8% vs 4%), lymphopenia (6% vs 3%), back pain (6% vs 3%), and pelvic pain (6% vs 1%)
- In mRCC, the most common Grade 3–5 adverse reactions in AVOREN, occurring at a >2% higher incidence in bevacizumab-treated patients vs controls, were fatigue (13% vs 8%), asthenia (10% vs 7%), proteinuria (7% vs 0%), hypertension (6% vs 1%, including hypertension and hypertensive crisis), and hemorrhage (3% vs 0.3%; including epistaxis, small intestinal hemorrhage, aneurysm ruptured, gastric ulcer hemorrhage, gingival bleeding, hemoptysis, hemorrhage intracranial, large intestinal hemorrhage, respiratory tract hemorrhage, and traumatic hematoma)
- In rGBM Study EORTC 26101, the incidence of Grade 3-4 VTE was 5% in patients receiving bevacizumab with chemotherapy compared to 2% in patients receiving chemotherapy alone. In this study, 22% of patients discontinued treatment

in the bevacizumab with lomustine arm due to adverse reactions compared with 10% of patients in the lomustine arm. In patients receiving



IMPORTANT SAFETY INFORMATION (cont'd)

bevacizumab with lomustine, the adverse reaction profile was similar to that observed in other approved indications

- In NSCLC, Grade 3–5 (nonhematologic) and Grade 4–5 (hematologic) adverse reactions in Study E4599 occurring at a ≥2% higher incidence in bevacizumab-treated patients vs controls were neutropenia (27% vs 17%), fatigue (16% vs 13%), hypertension (8% vs 0.7%), infection without neutropenia (7% vs 3%), venous thromboembolism (5% vs 3%), febrile neutropenia (5% vs 2%), pneumonitis/pulmonary infiltrates (5% vs 3%), infection with Grade 3 or 4 neutropenia (4% vs 2%), hyponatremia (4% vs 1%), headache (3% vs 1%), and proteinuria (3% vs 0%)
- In first-line MCRC, the most common Grade 3-4 reactions in Study 2107, which occurred at a ≥2% higher incidence in the bevacizumab plus IFL vs IFL groups, were asthenia (10% vs 7%), abdominal pain (8% vs 5%), pain (8% vs 5%), hypertension (12% vs 2%), deep vein thrombosis (9% vs 5%), intra-abdominal thrombosis (3% vs 1%), syncope (3% vs 1%), diarrhea (34% vs 25%), constipation (4% vs 2%), leukopenia (37% vs 31%), and neutropenia (21% vs 14%)
- In second-line MCRC, the most common Grade 3-5 (nonhematologic) and 4-5 (hematologic) reactions in Study E3200, which occurred at a higher incidence (≥2%) in the bevacizumab plus FOLFOX4 vs FOLFOX4 groups, were fatigue (19% vs 13%), diarrhea (18% vs 13%), sensory neuropathy (17% vs 9%), nausea (12% vs 5%), vomiting (11% vs 4%), dehydration (10% vs 5%), hypertension (9% vs 2%), abdominal pain (8% vs 5%), hemorrhage (5% vs 1%), other neurological (5% vs 3%), ileus (4% vs 1%), and headache (3% vs 0%). These data are likely to underestimate the true adverse event rates due to the reporting mechanisms used in this study
- When continued beyond first progression in MCRC, no new safety signals were observed in the TML study (ML18147) when bevacizumab was administered in second-line MCRC patients who progressed on a bevacizumab containing regimen

in first-line MCRC. The safety data was consistent with the known safety profile established in firstand second-line MCRC

You may report side effects to the FDA at (800) FDA-1088 or **www.fda.gov/medwatch.** You may also report side effects to Amgen at 1-800-772-6436.

Please see full Prescribing Information for additional important safety information.

INDICATIONS

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MVASI® is indicated for the treatment of recurrent glioblastoma (GBM) in adults.

MVASI[®], in combination with interferon-alfa, is indicated for the treatment of metastatic renal cell carcinoma (mRCC).

MVASI®, in combination with paclitaxel and cisplatin or paclitaxel and topotecan, is indicated for the treatment of patients with persistent, recurrent, or metastatic cervical cancer (CC).



SUPPORT SERVICES



See How We Can Help Your Patients

Offering the tools, information, and support for Amgen products that make a difference for you and your patients



AMGEN REIMBURSEMENT SPECIALISTS

Connect with an Amgen Reimbursement Counselor or schedule a visit with a Field Reimbursement Specialist



PATIENT RESOURCE GUIDE

Find co-pay and reimbursement resources* for patients with different kinds of insurance, or no insurance at all



BENEFIT VERIFICATION

Submit, store, and retrieve benefit verifications for all your patients currently on an Amgen product

*Resources include referrals to independent nonprofit patient assistance programs. Eligibility for resources provided by independent nonprofit patient assistance programs is based on the nonprofits' criteria. Amgen has no control over these programs and provides referrals as a courtesy only.

CALL **1-888-4ASSIST** (888-427-7478) Monday to Friday, 9:00 am to 8:00 pm ET, or visit **AmgenAssist360.com**.



NOTES





References: 1. MVASI[®] (bevacizumab-awwb) Prescribing Information, Amgen.
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6. American Medical Association. 2017 Professional Edition, Current Procedural Terminology (CPT) copyright 2016 American Medical Association. All rights reserved.

Reimbursement Disclaimer

This resource intended as a reference for coding and billing for product and associated services. It is not intended to be directive; the use of the recommended codes does not guarantee reimbursement. Healthcare providers may deem other codes or policies more appropriate and should select the coding options that most accurately reflect their internal system guidelines, payer requirements, practice patterns, and the services rendered. Healthcare providers are responsible for ensuring the accuracy and validity of all billing and claims for appropriate reimbursement.

Please see **full Important Safety Information** and **click here for full Prescribing Information**.

Please visit MVASI.com for additional information and resources.

Call **1-800-77-AMGEN (1-800-772-6436)** if you have questions about the preparation and administration of MVASI[®].

