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**UNITED STATES  
SECURITIES AND EXCHANGE COMMISSION  
WASHINGTON, D.C. 20549**

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**FORM 6-K**

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**REPORT OF FOREIGN PRIVATE ISSUER  
PURSUANT TO RULE 13a-16 OR 15d-16  
UNDER THE SECURITIES EXCHANGE ACT OF 1934**

For the Month of May 2026

Commission File Number: 001-38097

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**ARGENX SE**

(Translation of registrant's name into English)

Laarderhoogtweg 25  
1101 EB Amsterdam, the Netherlands  
(Address of principal executive offices)

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Indicate by check mark whether the registrant files or will file annual reports under cover of Form 20-F or Form 40-F.

Form 20-F  Form 40-F

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1):

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7):

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**EXPLANATORY NOTE**

On May 7, 2026, argenx SE (the "Company") issued a press release and an investor presentation, copies of which are attached hereto as Exhibits 99.1 and 99.2, respectively, and are incorporated by reference herein.

*The information contained in this Current Report on Form 6-K, including Exhibits 99.1 and 99.2, shall be deemed to be incorporated by reference into the Company's Registration Statements on Form S-8 (File Nos. [333-225375](#), [333-258253](#), [333-274721](#), and [333-292200](#)), and to be part thereof from the date on which this Current Report on Form 6-K is filed, to the extent not superseded by documents or reports subsequently filed or furnished.*

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**EXHIBITS**

<u>Exhibit</u>	<u>Description</u>
99.1	<a href="#">Press Release dated May 7, 2026</a>
99.2	<a href="#">Investor Presentation dated May 7, 2026</a>

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**SIGNATURES**

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

Date: May 7, 2026

**ARGENX SE**  
By: /s/ Hemamalini (Malini) Moorthy  
Hemamalini (Malini) Moorthy  
General Counsel

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## argenx Reports First Quarter 2026 Financial Results and Provides Business Update

*\$1.3 billion in first quarter global product net sales, representing 63% year-over-year growth*

*Anti-AChR antibody negative ("seronegative") gMG PDUFA is May 10, 2026*

*Management to host conference call today at 2:30 PM CET (8:30 AM ET)*

**May 7, 2026, 7:00AM CET**

**Amsterdam, the Netherlands** – argenx SE (Euronext & Nasdaq: ARGX), a global immunology company committed to improving the lives of people suffering from severe autoimmune diseases, today announced its first quarter 2026 results and provided a business update.

"argenx continues to deliver meaningful impact for patients, reflected by our 17th consecutive quarter of VYVGART growth," said Karen Massey, Chief Executive Officer of argenx. "Looking ahead, VYVGART has the potential to become the first and only approved therapy across MG, pending FDA decisions on label expansions into seronegative and ocular populations. At the same time, we are extending our leadership in FcRn into rheumatology, beginning with the upcoming myositis readout. Our next pipeline candidate, empasiprubart, is progressing toward its first registrational readout in MMN, and we continue to advance a broad and differentiated pipeline. With these opportunities, we remain focused on delivering transformative outcomes for patients while creating sustained value for all stakeholders."

### **Vision 2030**

argenx continues to advance its 'Vision 2030' anchored in the ambition to treat 50,000 patients globally with its medicines, secure 10 labeled indications, and progress five pipeline candidates into Phase 3 development by 2030.

### **Expanding global VYVGART opportunity and shaping the long-term future of FcRn**

VYVGART® (IV: efgartigimod alfa-fcab and SC: efgartigimod alfa and hyaluronidase-qvfc) is a first-and-only IgG Fc-antibody fragment that targets the neonatal Fc receptor (FcRn). It is approved in three indications, including generalized myasthenia gravis (gMG) and chronic inflammatory demyelinating polyneuropathy (CIDP) globally, and primary immune thrombocytopenia (ITP) in Japan. argenx is driving broad adoption as the leading precision biologic in MG and CIDP while advancing multiple label expansions. argenx is also shaping the future of FcRn medicines by advancing new pipeline candidates and delivery modalities.

- Generated \$1.3 billion in global product net sales in the first quarter of 2026, representing an increase of approximately 63% or \$0.5 billion in year-over-year growth
- Prescription Drug User Fee Act (PDUFA) target action date for anti-acetylcholine receptor antibody negative (AChR-Ab-) gMG (MuSK+, LRP4+ and triple seronegative) is May 10, 2026
- Positive topline results from ADAPT OCULUS were recently presented at AAN; these data support planned sBLA submission to expand VYVGART label into oMG
- Topline results from ALKIVIA study (myositis) expected in third quarter of 2026
- Topline results from ADVANCE-NEXT study (primary ITP) expected in first half of 2027
- Registrational study in Graves' disease (GD) expected to initiate in 2026, expanding development into thyroid-driven autoimmunity
- Topline results from UNITY study (Sjogren's disease) expected in second half of 2027
- VYVGART SC autoinjector expected to launch in 2027 for all approved indications
- Progressing two future FcRn molecules: ARGX-213 is Phase 3-ready and ARGX-124 is in Phase 1

**Advancing empasiprubart**

Empasiprubart is a first-in-class, humanized monoclonal antibody designed to inhibit complement factor C2, selectively blocking activation of the classical and lectin complement pathways. It is being evaluated in registrational studies in multifocal motor neuropathy (MMN) and CIDP, and in a combination study with VYVGART in gMG.

- Topline results from EMPASSION study (MMN) expected in fourth quarter of 2026
- Topline results from EMVIGORATE and EMNERGIZE studies (CIDP) expected in second half of 2027
- Decision for Phase 2 VARVARA study (Delayed Graft Function) expected mid-year 2026 following completion of 52-week efficacy analysis
- ADAPT-Forward combination study ongoing to evaluate empasiprubart as an add on therapy to efgartigimod in gMG

**Delivering next wave of immunology innovation**

By the end of 2026, the argenx pipeline is expected to include a total of ten molecules in clinical development. Beyond efgartigimod and empasiprubart, this includes adimanebart (a MuSK agonist); ARGX-121 (anti-IgA), ARGX-109 (anti-IL-6), and three additional molecules from the Immunology Innovation Program (IIP). Collectively, these programs support argenx's goal of launching, on average, one new pipeline candidate per year.

- Adimanebart CMS registrational study on track to start in third quarter of 2026
- Phase 2 study of ARGX-121 in IgA nephropathy (IgAN) expected to start in 2026
- Three new first-in-class molecules on track to enter Phase 1 in 2026, including ARGX-118 (Galectin-10 inhibitor), ARGX-125 (bispecific antibody), and TSP-101, the Fn14-targeting program from the Tensegrity research collaboration

**Key business highlights**

- On May 6, 2026, Karen Massey was appointed Chief Executive Officer and executive director of the argenx Board of Directors following the Annual General Meeting of Shareholders. Tim Van Hauwermeiren was appointed non-executive director and Chairperson of the Board of Directors
- In March 2026, argenx expanded its global presence in Asia with the establishment of an argenx affiliate in China to broaden its access to novel biology and support early-stage research

FIRST QUARTER 2026 FINANCIAL RESULTS

argenx SE

UNAUDITED CONDENSED CONSOLIDATED INTERIM STATEMENTS OF PROFIT OR LOSS

(in millions of \$ except for per share data)	Three Months Ended	
	March 31,	
	2026	2025
Product net sales	\$ 1,298	\$ 790
Other operating income*	15	17
<b>Total operating income</b>	<b>1,313</b>	<b>807</b>
Cost of sales	\$ (121)	\$ (81)
Research and development expenses*	(443)	(311)
Selling, general and administrative expenses	(355)	(276)
<b>Total operating expenses</b>	<b>(919)</b>	<b>(668)</b>
<b>Operating profit</b>	<b>\$ 394</b>	<b>\$ 139</b>
Financial income	\$ 44	\$ 37
Financial expense	(1)	(1)
Exchange (losses)/gains	(11)	27
<b>Profit for the period before taxes</b>	<b>\$ 426</b>	<b>\$ 202</b>
Income tax expense	\$ (60)	\$ (33)
<b>Profit for the period</b>	<b>\$ 366</b>	<b>\$ 169</b>
Profit for the period attributable to:		
Owners of the parent	\$ 366	\$ 169
Weighted average number of shares outstanding	62,056,886	60,983,325
Basic profit per share (in \$)	\$ 5.90	\$ 2.78
Weighted average number of shares outstanding for diluted profit per share	66,356,591	65,664,300
Diluted profit per share (in \$)	\$ 5.52	\$ 2.58

\*Comparative figures have been aligned with the presentation adopted in the current period, reflecting the combination of collaboration revenue and other operating income, as well as the combination of research and development expenses and loss from investment in a joint venture.

## DETAILS OF THE FINANCIAL RESULTS

**Total operating income** for the three months ended March 31, 2026, was \$1.3 billion compared to \$0.8 billion for the same period in 2025, and consists of:

- **Product net sales** of VYVGART for the three months ended March 31, 2026, were \$1.3 billion compared to \$0.8 billion for the same period in 2025.
- **Other operating income** for the three months ended March 31, 2026, was \$15 million compared to \$17 million for the same period in 2025. The other operating income primarily relates to research and development tax incentives and payroll tax rebates.

**Total operating expenses** for the three months ended March 31, 2026, were \$0.9 billion compared to \$0.7 billion for the same period in 2025, and mainly consists of:

- **Cost of sales** for the three months ended March 31, 2026, was \$121 million compared to \$81 million for the same period in 2025. The cost of sales was recognized with respect to the sale of VYVGART.
- **Research and development expenses** for the three months ended March 31, 2026, were \$0.4 billion compared to \$0.3 billion for the same period in 2025. The expenses mainly relate to:
  - Advancing efgartigimod across multiple severe autoimmune diseases;
  - Progressing empasiprubar into multiple indications;
  - Executing studies for adimanebart in rare neuromuscular diseases; and
  - Early-stage discovery and preclinical programs to sustain long-term pipeline growth.
- **Selling, general and administrative expenses** for the three months ended March 31, 2026, were \$0.4 billion compared to \$0.3 billion for the same period in 2025. The selling, general and administrative expenses mainly relate to professional and marketing fees linked to global commercialization of the VYVGART franchise, and personnel expenses.

**Financial income** for the three months ended March 31, 2026, was \$44 million compared to \$37 million for the same period in 2025.

**Income tax expense** for the three months ended March 31, 2026 was \$60 million compared to \$33 million for the same period in 2025. Income tax expense for the three months ended March 31, 2026, consists of \$102 million of current income tax expense and \$42 million of deferred tax benefit, compared to \$29 million of current income tax expense and \$4 million of deferred tax expense for the comparable prior period.

**Profit** for the period of three months ended March 31, 2026, was \$366 million compared to \$169 million in 2025, representing 116% growth year-over-year. The basic profit per share was \$5.90 for the three months ended March 31, 2026 compared to \$2.78 in 2025.

**Cash, cash equivalents and current financial assets**<sup>1</sup> consisted of \$4.3 billion in cash, cash equivalents and \$0.6 billion in current financial assets which totaled \$4.9 billion as of March 31, 2026, compared to \$3.5 billion in cash and cash equivalents and \$0.9 billion in current financial assets which totaled \$4.4 billion as of December 31, 2025.

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<sup>1</sup> A non-IFRS Alternative Performance Measure (APM). Refer to the "Alternative Performance Measures Statement" below for a reconciliation to the IFRS financial information.



## EXPECTED 2026 FINANCIAL CALENDAR

- July 23, 2026: Half Year and Second Quarter 2026 Financial Results and Business Update
- October 22, 2026: Third Quarter 2026 Financial Results and Business Update

## CONFERENCE CALL DETAILS

The first quarter 2026 financial results and business update will be discussed during a conference call and webcast presentation today at 2:30 PM CET/8:30 AM ET. A webcast of the live call may be accessed on the Investors section of the argenx website at [argenx.com/investors](https://argenx.com/investors). A replay of the webcast will be available on the argenx website.

### Dial-in numbers:

*Please dial in 15 minutes prior to the live call.*

Belgium	32 800 50 201
France	33 800 943355
Netherlands	31 20 795 1090
United Kingdom	44 800 358 0970
United States	1 888 415 4250
Japan	81 3 4578 9081
Switzerland	41 43 210 11 32

## About VYVGART

VYVGART® (efgartigimod alfa fcab) is a human IgG1 antibody fragment that binds to the neonatal Fc receptor (FcRn), resulting in the reduction of circulating IgG autoantibodies. It is the first approved FcRn blocker for the treatment of generalized myasthenia gravis (gMG) and chronic inflammatory demyelinating polyneuropathy (CIDP) globally, and for primary immune thrombocytopenia (ITP) in Japan. VYVGART SC is a subcutaneous combination of efgartigimod alfa and recombinant human hyaluronidase PH20 (rHuPH20), Halozyme's ENHANZE® drug delivery technology to facilitate subcutaneous injection delivery of biologics. It is marketed as VYVGART® Hytrulo in the U.S., VYVGART SC in Europe, VYVDURA® in Japan, and may be marketed under different proprietary names following approval in other regions.

## About argenx

argenx is a global immunology company committed to improving the lives of people suffering from severe autoimmune diseases. Partnering with leading academic researchers through its Immunology Innovation Program (IIP), argenx aims to translate immunology breakthroughs into a world-class portfolio of novel antibody-based medicines. argenx developed and is commercializing the first approved neonatal Fc receptor (FcRn) blocker and is evaluating its broad potential in multiple serious autoimmune diseases while advancing several earlier stage experimental medicines within its therapeutic franchises. For more information, visit [www.argenx.com](https://www.argenx.com) and follow us on [LinkedIn](#), [Instagram](#), [Facebook](#), and [YouTube](#).

**This press release contains inside information within the meaning of Article 7(1) of the EU Market Abuse Regulation (Regulation 596/2014).**

**For further information, please contact:**

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[aroy@argenx.com](mailto:aroy@argenx.com)

## Forward-looking Statements

The contents of this announcement include statements that are, or may be deemed to be, "forward-looking statements." These forward-looking statements can be identified by the use of forward-looking terminology, including the terms "advance," "aim," "commit," "continue," "drive," "is," "potential," "reinforce," "represent," and



"will," and include statements argenx makes concerning its belief in VYVGART's potential to become the first and only approved therapy across MG, pending FDA decisions on label expansions into seronegative and ocular populations; its extension of its leadership in FcRn into rheumatology, beginning with myositis; the progression of its next pipeline candidate, empasiprubart, towards its first registrational readout in MMN; its advancement of a broad and differentiated pipeline; its focus on delivering transformative outcomes for patients while creating sustained value for all stakeholders; its advancement of its "Vision 2030" anchored in the ambition to treat 50,000 patients globally with its medicines, secure 10 labeled indications, and progress five pipeline candidates into Phase 3 development by 2030; driving broad adoption as the leading precision biologic in MG and CIDP while advancing multiple label expansions; its belief that it is also shaping the future of FcRn medicines by advancing new pipeline candidates and delivery modalities; the Prescription Drug User Fee Act (PDUFA) target action date of May 10, 2026 for anti-acetylcholine receptor antibody negative (AChR-Ab-) gMG (MuSK+, LRP4+ and triple seronegative); its planned sBLA submission to expand VYVGART label into oMG; its topline results from ALKIVIA study (myositis) expected in third quarter of 2026; its topline results expected for primary ITP (ADVANCE-NEXT) in the first half of 2027; its registrational study in Graves' disease (GD) expected to initiate in 2026, expanding development into thyroid-driven autoimmunity; its topline results from UNITY study (Sjogren's disease) expected in second half of 2027; its VYVGART SC autoinjector expected to launch in 2027 for all approved indications; its progression of two future FcRn molecules in 2026: ARGX-213 expected to enter patient studies, and ARGX-124 expected to complete Phase 1 development; its advancement of empasiprubart, including (1) topline results from EMPASSION study (MMN) expected in fourth quarter of 2026; (2) topline results from EMVIGORATE and EMNERGIZE studies (CIDP) expected in second half of 2027; (3) the decision for Phase 2 VARVARA study (Delayed Graft Function, DGF) expected mid-year 2026 following completion of 52-week efficacy analysis; and (4) the ADAPT-Forward combination study ongoing to evaluate empasiprubart as an add on therapy to efgartigimod in gMG; its expectation that the argenx pipeline will include a total of ten molecules in clinical development, including: adimanebart (a MuSK agonist), which is expected to enter Phase 3 development in congenital myasthenic syndromes (CMS); ARGX-121 (anti-IgA) and ARGX-109 (anti-IL-6), both of which are advancing into Phase 2 studies; and three additional molecules from the Immunology Innovation Program (IIP) on track to enter Phase 1 in 2026; its belief that these programs collectively support its goal of launching, on average, one new pipeline candidate per year; and its belief that (1) the CMS registrational study for Adimanebart is on track to start in third quarter of 2026; (2) the Phase 2 study of ARGX-121 in IgA nephropathy (IgAN) is expected to start in 2026; and (3) three new first-in-class molecules are on track to enter Phase 1 in 2026, including ARGX-118 (Galectin-10 inhibitor), ARGX-125 (bispecific antibody), and TSP-101, the Fn14-targeting program from the Tensegry research collaboration. By their nature, forward-looking statements involve risks and uncertainties and readers are cautioned that any such forward-looking statements are not guarantees of future performance. argenx's actual results may differ materially from those predicted by the forward-looking statements as a result of various important factors, including but not limited to, the results of argenx's clinical trials; expectations regarding the inherent uncertainties associated with the development of novel drug therapies; preclinical and clinical trial and product development activities and regulatory approval requirements; the acceptance of its products and product candidates by its patients as safe, effective and cost-effective; the impact of governmental laws and regulations, including tariffs, export controls, sanctions and other regulations on its business; its reliance on third-party suppliers, service providers and manufacturers; inflation and deflation and the corresponding fluctuations in interest rates; and regional instability and conflicts. A further list and description of these risks, uncertainties and other risks can be found in argenx's U.S. Securities and Exchange Commission (SEC) filings and reports, including in argenx's most recent annual report on Form 20-F filed with the SEC as well as subsequent filings and reports filed by argenx with the SEC. Given these uncertainties, the reader is advised not to place any undue reliance on such forward-looking statements. These forward-looking statements speak only as of the date of publication of this document. argenx undertakes no obligation to publicly update or revise the information in this press release, including any forward-looking statements, except as may be required by law.

#### **Alternative Performance Measures Statement**

In this document, argenx's financial results are provided in accordance with IFRS® Accounting Standards (IFRS) and using a non-IFRS financial measure, cash, cash equivalents and current financial assets.

This value should not be viewed as a substitute for the company's IFRS financial information and is provided as a complement to financial information provided in accordance with IFRS and should be read in conjunction with the most directly comparable IFRS financial information as set out below. Management believes this non-IFRS financial measure is useful for securities analysts, investors and other interested parties to gain a more complete understanding of the company's available financial liquidities given that the company's current financial assets are held in term accounts with an initial maturity of more than three months but less than twelve that may be used

to meet its financial obligations. Such non-IFRS financial information, as calculated herein, may not be comparable to similarly named measures used by other companies and should not be considered comparable to IFRS financial measures. Non-IFRS financial measures have limitations as an analytical tool and should not be considered in isolation from, or as a substitute for, an analysis of the company's financial results as reported under IFRS.

A reconciliation of the IFRS financial information to non-IFRS financial information is included below:

Cash, cash equivalents and current financial assets totaled \$4.9 billion as of March 31, 2026, compared to \$4.4 billion as of December 31, 2025. The balance as of the period ended March 31, 2026 consisted of \$4.3 billion in cash, cash equivalents and \$0.6 billion in current financial assets and the balance as of the period ended December 31, 2025 consisted of \$3.5 billion in cash and cash equivalents and \$0.9 billion in current financial assets.

Leading a new era of  
innovation in immunology

1Q 2026 FINANCIAL RESULTS CALL  
MAY 7, 2026



# Forward Looking Statements

This presentation has been prepared by argenx se ("argenx" or the "company") for informational purposes only and not for any other purpose. Nothing contained in this presentation is, or should be construed as, a recommendation, promise or representation by the presenter or the company or any director, employee, agent, or adviser of the company. This presentation does not purport to be all-inclusive or to contain all of the information you may desire. Certain information contained in this presentation relates to or is based on studies, publications, surveys and other data obtained from third-party sources and the company's own internal estimates and research. While argenx believes these third-party studies, publications, surveys and other data to be reliable as of the date of this presentation, it has not independently verified, and makes no representation as to the adequacy, fairness, accuracy or completeness of, any information obtained from third-party sources. In addition, no independent source has evaluated the reasonableness or accuracy of argenx's internal estimates or research, and no reliance should be made on any information or statements made in this presentation relating to or based on such internal estimates and research.

Certain statements contained in this presentation, other than present and historical facts and conditions independently verifiable at the date hereof, may constitute forward-looking statements. These forward-looking statements can be identified by the use of forward-looking terminology, including the terms "bring," "build," "commit," "continue," "expand," "expect," "grow," "is," "potential," and "will," and include statements argenx makes regarding its commitment to its transformation mission, Vision 2030, with 5 new molecules in Phase 3, 10 labeled indications, and 50,000 patients on treatment; its transformative potential across pipeline programs, including Complement Factor C2, C2-Specific Antibody, and Empasiprubarat, with Empasiprubarat's MMN registration readout expected in 4Q 2026 and CIDP registration readout expected in the second half of 2027; its commitment to building its presence in rheumatology by redefining biology, treatment, and patient outcomes in Autoimmune Myositis and the potential to reduce steroid burden and reliance on other conventional treatments; its transformative potential across pipeline programs (Elgartigimod, Empasiprubarat, Adimanebart, ARGX-213 and ARGX-121); and its ability to sustain and grow MG and CIDP markets and leadership.

By their nature, forward-looking statements involve risks and uncertainties and readers are cautioned that any such forward-looking statements are not guarantees of future performance. argenx's actual results may differ materially from those predicted by the forward-looking statements as a result of various important factors, including the results of argenx's clinical trials; expectations regarding the inherent uncertainties associated with the development of novel drug therapies; preclinical and clinical trial and product development activities and regulatory approval requirements in products and product candidates; the acceptance of argenx's products and product candidates by patients as safe, effective and cost-effective; the impact of governmental laws and regulations on our business, including tariffs, export controls, sanctions and other regulations on its business; disruptions caused on our reliance on third parties suppliers, service providers and manufacturing; inflation and deflation and the corresponding fluctuations in interest rates; and regional instability and conflicts. A further list and description of these risks, uncertainties and other risks can be found in argenx's U.S. Securities and Exchange Commission (the "SEC") filings and reports, including in argenx's most recent annual report on Form 20-F filed with the SEC as well as subsequent filings and reports filed by argenx with the SEC. Given these uncertainties, the reader is advised not to place any undue reliance on such forward-looking statements. These forward-looking statements speak only as of the date of publication of this document. argenx undertakes no obligation to publicly update or revise the information in this presentation, including any forward-looking statements, except as may be required by law.

This presentation contains trademarks, trade names and service marks of other companies, which are the property of their respective owners.

## Commitment To Our Transformation Mission

Disciplined Scaling

Leadership in FcRn

Continuous Pipeline of Innovation

# VISION 2030

**50k** Patients on Treatment

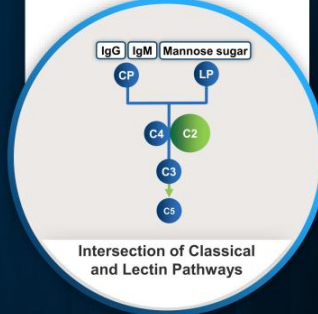
**10** Labeled Indications

**5** New Molecules in Phase 3

# Empasiprubart: Our Second Potential Medicine

Foundational  
Immune Target

## Complement Factor C2



Intersection of Classical  
and Lectin Pathways

First-in-Class  
Potential Best-in-Class

## C2-Specific Antibody



NHance™

Pipeline in a  
Product Opportunity

## Empasiprubart

**MMN**  
Registrational Readout  
4Q 2026

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**CIDP**  
Registrational Readout  
2H 2027

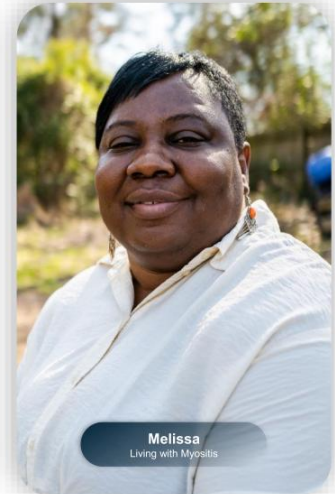
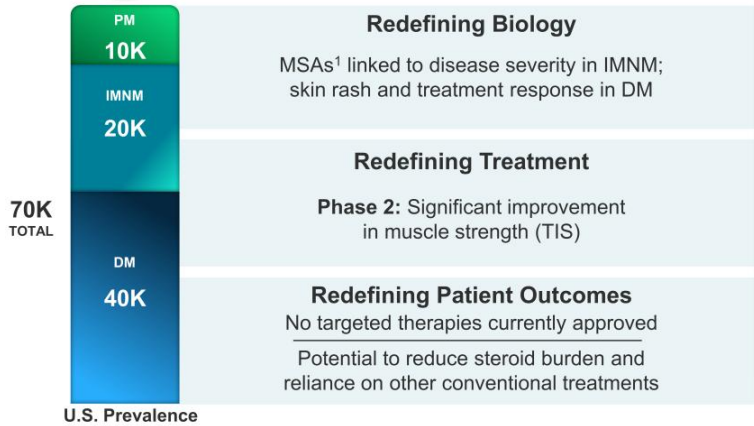
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Explored in Combination

# Building our Presence in Rheumatology



## Autoimmune Myositis



**Melissa**  
Living with Myositis

1.MSA: myositis specific antibodies (anti-SRP and -HMGR in IMNM, anti-Mi2 and -MDA5 in DM)

# Transformative Potential Across Pipeline Programs

Efgartigimod



First-in-class  
Fc Fragment

**15+**

Indications

Empasiprubart



Potent C2  
sweeping antibody

**3+**

Indications

Adimanebart



MuSK agonist  
antibody

**3+**

Indications

ARGX-213



FcRn: Sustained  
IgG reduction

**15+**

*Potential* indications

ARGX-121



IgA Sweeping  
Antibody

**3+**

*Potential* indications

# Product Net Sales of \$1.3 Billion in Q1



**Year-over-Year Growth of 63%\***

**Q1 2026 growth vs Q1 2025**

(in millions of \$)

	Q1 2026	Q1 2025	Growth	Growth % *
US	1,107	681	426	62%
Japan	67	32	35	111%
Rest of the World	112	57	55	99%
China supply	12	20	(8)	(41%)
<b>Total</b>	<b>1,298</b>	<b>790</b>	<b>508</b>	<b>63%</b>

**Q1 2026 growth vs Q4 2025**

(in millions of \$)

	Q1 2026	Q4 2025	Growth	QoQ % Growth *
US	1,107	1,087	20	2%
Japan	67	63	4	6%
Rest of the World	112	110	2	3%
China supply	12	26	(14)	(54%)
<b>Total</b>	<b>1,298</b>	<b>1,286</b>	<b>12</b>	<b>1%</b>
<b>Total ex-China</b>	<b>1,286</b>	<b>1,260</b>	<b>26</b>	<b>2%</b>

\*Product Net sales growth % excludes the impact of FX

**VVVGART™**  
(efgartigimod alfa-fcab)

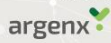
**VVVGART Hytrulo**  
(efgartigimod alfa and hyaluronidase-qvfc)  
Subcutaneous injection, 90 mg/mL and 2000 U/mL, vial

argenx 

# Q1 2026 Financial Summary

(in million of \$)	Three months ended	
	March 31	
	2026	2025
Product net sales	1,298	790
Other operating income*	15	17
<b>Total operating income</b>	<b>1,313</b>	<b>807</b>
Cost of sales	(121)	(81)
Research and development expenses*	(443)	(311)
Selling, general and administrative expenses	(355)	(276)
<b>Total operating expenses</b>	<b>(919)</b>	<b>(668)</b>
<b>Operating profit</b>	<b>394</b>	<b>139</b>
Financial income	44	37
Financial expense	(1)	(1)
Exchange (losses)/gains	(11)	27
<b>Profit for the period before taxes</b>	<b>426</b>	<b>202</b>
Income tax expense	(60)	(33)
<b>Profit for the period</b>	<b>366</b>	<b>169</b>

\*Comparative figures have been aligned with the presentation adopted in the current period, reflecting the combination of collaboration revenue and other operating income, as well as the combination of research and development expenses and loss from investment in a joint venture.



**Delivering Margin Expansion**  
**Operating profit of \$394 million, +183% YoY**

**\$4.3 billion in cash and cash equivalents and \$0.6 billion in current financial assets**  
**Ended Q1 with cash<sup>†</sup> of \$4.9B**

<sup>†</sup> Alternative Performance Measure (APM). Refer to the APM Statement.

# Raising the Standard of Care for Patients



Securing broad and simple patient access



Driving earlier-line use of VYVGART in MG and CIDP



Disciplined, scalable commercial execution

Jai, VYVGART Patient

## Strong Momentum Across MG and CIDP



Mary Beth, MG Patient

### NEW PATIENT STARTS

Amongst  
**Highest Achieved**  
since launch

### PRESCRIBER EXPANSION

**~5,000**  
Prescribers in the US  
2X increase since CIDP launch

### EARLIER LINE USE

**4/5**  
Prescribers in the US  
Prefer to start with VYVGART as  
the targeted biologic in gMG

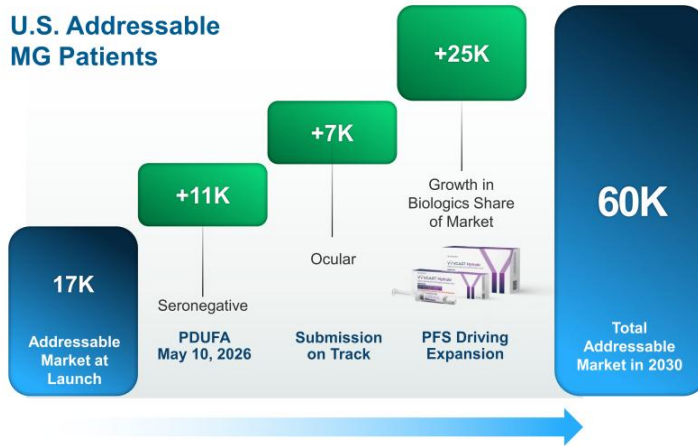
### PFS EXPANDING DEMAND

**68%**  
of PFS patients  
new to VYVGART since launch

Source: argenx market research  
As of 1Q 2026 Financial Results

# Sustaining MG Leadership

## U.S. Addressable MG Patients



*"I can carry a bag of potting soil from my car to the patio. I could (and wanted to!) work in my flower beds for longer than I'd been able to prior to treatment with VYVGART."*

**Pam**  
gMG Patient



# Raising the Bar in CIDP

Early-line use



## 87.5%

Clinical responses *observed among treatment-naïve patients* in ADHERE post hoc analysis

Sustained functional benefit



## 96 wks

Mean grip strength *continued to improve up to 96 weeks* in open-label extension



Scott, CIDP Patient

AAN Presentation: ADHERE Study post-hoc analysis from open label Stage A  
Lewis et. al: Impact of Efgartigimod PH20 SC on Grip Strength in CIDP: Post Hoc Analysis of ADHERE/ADHERE+

argenx

# Innovation Has No Value Unless It Provides Meaningful Benefit to Patients



## Alternative Performance Measure Statement

In this document, argenx's financial results are provided in accordance with IFRS® Accounting Standards (IFRS) and using a non-IFRS financial measure, cash, cash equivalents and current financial assets.

This value should not be viewed as a substitute for the company's IFRS financial information and is provided as a complement to financial information provided in accordance with IFRS and should be read in conjunction with the most directly comparable IFRS financial information as set out below. Management believes this non-IFRS financial measure is useful for securities analysts, investors and other interested parties to gain a more complete understanding of the company's available financial liquidities given that the company's current financial assets are held in term accounts with an initial maturity of more than three months but less than twelve that may be used to meet its financial obligations. Such non-IFRS financial information, as calculated herein, may not be comparable to similarly named measures used by other companies and should not be considered comparable to IFRS financial measures. Non-IFRS financial measures have limitations as an analytical tool and should not be considered in isolation from, or as a substitute for, an analysis of the company's financial results as reported under IFRS.

A reconciliation of the IFRS financial information to non-IFRS financial information is included below:

Cash, cash equivalents and current financial assets totaled \$4.9 billion as of March 31, 2026, compared to \$4.4 billion as of December 31, 2025. The balance as of the period ended March 31, 2026 consisted of \$4.3 billion in cash, cash equivalents and \$0.6 billion in current financial assets and the balance as of the period ended December 31, 2025 consisted of \$3.5 billion in cash and cash equivalents and \$0.9 billion in current financial assets.

