UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

FORM 8-K

CURRENT REPORT Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): October 7, 2019

(Exac	t name of registrant as specified in its	charter)
Delaware	001-36728	56-2590442
(State or other jurisdiction	(Commission	(IRS Employer
of incorporation)	File Number)	Identification No.)
465 State Route 17, Ramsey, New Jersey		07446
(Address of principal executive offices)		(Zip Code)
Registrant's to	elephone number, including area code:	z (<u>201) 478-5552</u>
(Former na	nme or former address, if changed sinc	re last report.)
Check the appropriate box below if the Form 8-K filing is a provisions (see General Instruction A.2. below):	intended to simultaneously satisfy the	filing obligation of the registrant under any of the following
o Written communications pursuant to Rule 425 under the	Securities Act (17 CFR 230.425)	
\square Soliciting material pursuant to Rule 14a-12 under the Ex	schange Act (17 CFR 240.14a-12)	
□ Jonething material pursuant to Rule 14a-12 under the L		
\square Pre-commencement communications pursuant to Rule 1	4d-2(b) under the Exchange Act (17 C	CFR 240.14d-2(b))
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☐ Pre-commencement communications pursuant to Rule 1	.,	· //
☐ Pre-commencement communications pursuant to Rule 1 ☐ Pre-commencement communications pursuant to Rule 1	.,	· //

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or

this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. \Box

Emerging growth company \square

Item 8.01 Other Events.

On October 7, 2019, ADMA Biologics, Inc. issued a press release entitled "ADMA Biologics Announces Data Presented at IDWeek 2019." The full text of the press release is filed as Exhibit 99.1 to this Current Report on Form 8-K and is incorporated herein by reference.

Item 9.01 Exhibits.

(d) Exhibits

Exhibit No. Description

99.1 Press release dated October 7, 2019, entitled "ADMA Biologics Announces Data Presented at IDWeek 2019."

SIGNATURE

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 hereunto duly authorized.

October 7, 2019 ADMA Biologics, Inc.

By: /s/ Brian Lenz

Name: Brian Lenz

Title: Executive Vice President and Chief

Financial Officer



ADMA Biologics Announces Data Presented at IDWeek 2019

RAMSEY, N.J. and BOCA RATON, FL., – October 7, 2019 – ADMA Biologics, Inc. (NASDAQ: ADMA) ("ADMA" or the "Company"), a vertically integrated commercial biopharmaceutical and specialty immunoglobulin company that manufactures, markets and develops specialty plasma-derived biologics for the treatment of immune deficiencies and the prevention of certain infectious diseases, announces that a poster presentation was given at IDWeek 2019, the combined annual meeting of IDSA, SHEA, HVMA and PIDS in Washington, D.C. The poster detailed data obtained from the compassionate use of ASCENIVTM (formerly referred to as RI-002) in the treatment of Respiratory Syncytial Virus ("RSV") infection in two immunocompromised children.

The two immunocompromised children admitted to the Mayo Clinic each were diagnosed with T-cell lymphoblastic lymphoma. Both patients were undergoing delayed intensification chemotherapy and each were diagnosed with RSV Lower Respiratory Tract Infection ("LRTI"). Both children were treated with ASCENIVTM under an emergency United States Food and Drug Administration ("FDA") Investigational New Drug protocol.

The treatment and results of patient one are as follows - Patient was admitted with fever, neutropenia, nasal congestion and diagnosed with RSV infection on hospital day five. On hospital day 17, the patient was intubated for respiratory failure. Intravenous Immune Globulin ("IVIG") and palivizumab, and daily oral ribavirin were administered. On hospital day eighteen, the patient required high frequency oscillator ventilation, nitric oxide and paralysis. The patient was then administered ASCENIVTM 1.5g/kg on hospital day twenty and 0.75g/kg on hospital day twenty-two. The patient was placed on veno-venous Extracorporeal Membrane Oxygenation ("ECMO") on hospital day twenty-three. On hospital day thirty-three, a third dose of ASCENIVTM was given at 0.75 g/kg. The patient's pulmonary compliance and chest CTs improved. On day fifty-two, ECMO support was discontinued and the patient was discharged from the hospital day eighty-eight, and currently requires no respiratory support.

The treatment and results of patient two are as follows - Patient was admitted with fever, neutropenia, nasal congestion, cough and stridor and diagnosed with RSV infection on hospital day one. The patient required nasal cannula oxygen. IVIG and daily oral ribavirin were administered. The patient was administered ASCENIVTM 1.5g/kg on hospital day three and 0.75g/kg on hospital day five. By hospital day five, the patient was afebrile; oxygen was discontinued and the patient was discharged from the hospital on hospital day six.

The data in the poster and abstract concludes that the evaluated product may be useful in the treatment of severe RSV infections and may assist in prevention of progression of RSV lower respiratory tract infection. The data also confirms that further evaluation of ASCENIVTM in this patient population is warranted.

About ADMA Biologics, Inc. (ADMA)

ADMA Biologics is a vertically integrated commercial biopharmaceutical company that manufactures and markets three United States Food and Drug Administration ("FDA") approved plasma-derived biologics for the treatment of immune deficiencies and the prevention of certain infectious diseases. ADMA's mission is to manufacture, market and develop plasma-derived, human immune globulins targeted to niche patient populations for the treatment and prevention of certain infectious diseases and management of immune compromised patient populations who suffer from an underlying immune deficiency disease, or who may be immune compromised for other medical reasons. ADMA has received U.S. Patents 9,107,906, 9,714,283, 9,815,886, 9,969,793 and 10,259,865 related to certain aspects of its products and product candidates. For more information, please visit www.admabiologics.com.

About ASCENIVTM (Formerly referred to as RI-002)

ASCENIV™, Immune Globulin Intravenous, Human – slra 10% Liquid, is a plasma-derived, polyclonal, intravenous immune globulin ("IVIG"). ASCENIV™ is protected by U.S. Patents: 9,107,906, 9,714,283 and 9,815,886. ASCENIV™ is manufactured using our unique, patented plasma donor screening methodology and tailored plasma pooling design, which blends normal source plasma and plasma from donors tested using our proprietary microneutralization assay. ASCENIV™ contains naturally occurring polyclonal antibodies. ASCENIV™ is indicated for the treatment of Primary Humoral Immunodeficiency or Primary Immune Deficiency Disease ("PI") in adults and adolescents (12 to 17 years of age). ADMA received FDA approval for ASCENIV™ on April 1, 2019. Polyclonal antibodies are proteins that are used by the body's immune system to neutralize microbes, such as bacteria and viruses and prevent against infection and disease. ASCENIV™ prevented serious bacterial infection among 59 patients treated for twelve months during the pivotal investigation. The most common adverse reactions to ASCENIV™ (≥5% of study subjects) were headache, sinusitis, diarrhea, gastroenteritis viral, nasopharyngitis, upper respiratory tract infection, bronchitis, and nausea. ADMA anticipates the commercial launch of ASCENIV™ during the fourth quarter of 2019. Certain data and other information about ASCENIV™ or ADMA Biologics and its products can be found on the Company's website at: www.admabiologics.com.

Additional Important Safety Information about ASCENIVTM

ASCENIV™ (immune globulin intravenous, human – slra) is a 10% immune globulin liquid for intravenous injection, indicated for the treatment of primary humoral immunodeficiency (PI) in adults and adolescents (12 to 17 years of age). PI includes, but is not limited to, the humoral immune defect in congenital agammaglobulinemia, common variable immunodeficiency (CVID), X linked agammaglobulinemia, Wiskott-Aldrich syndrome, and severe combined immunodeficiencies (SCID).

WARNING: THROMBOSIS, RENAL DYSFUNCTION AND ACUTE RENAL FAILURE

Thrombosis may occur with immune globulin (IGIV) products, including ASCENIV TM . Risk factors may include: advanced age, prolonged immobilization, hypercoagulable conditions, history of venous or arterial thrombosis, use of estrogens, indwelling central vascular catheters, hyperviscosity, and cardiovascular risk factors. Thrombosis may occur in the absence of known risk factors.

Renal dysfunction, acute renal failure, osmotic nephrosis, and death may occur with the administration of IGIV products in predisposed patients.

Renal dysfunction and acute renal failure occur more commonly in patients receiving IGIV products containing sucrose. ASCENIV TM does not contain sucrose.

For patients at risk of thrombosis, renal dysfunction or renal failure, administer $ASCENIV^{TM}$ at the minimum dose and infusion rate practicable. Ensure adequate hydration in patients before administration. Monitor for signs and symptoms of thrombosis and assess blood viscosity in patients at risk for hyperviscosity.

ASCENIVTM is contraindicated in:

- Patients who have had an anaphylactic or severe systemic reaction to the administration of human immune globulin.
- · IgA-deficiency patients with antibodies to IgA and a history of hypersensitivity.

Warnings and Precautions

Severe hypersensitivity reactions may occur with IGIV products, including ASCENIVTM. In case of hypersensitivity, discontinue ASCENIVTM infusion immediately and institute appropriate treatment. Medications such as epinephrine should be available for treatment of acute hypersensitivity reactions.

Thrombosis may occur following treatment with immunoglobulin products, including ASCENIVTM. Thrombosis may occur in the absence of known risk factors.

Acute renal dysfunction/failure, osmotic nephrosis, and death may occur upon use of human IGIV products. Ensure that patients are not volume depleted before administering ASCENIV™. Periodic monitoring of renal function and urine output is particularly important in patients judged to be at increased risk of developing acute renal failure.

Hyperproteinemia, increased serum viscosity, and hyponatremia may occur in patients receiving IGIV treatment, including ASCENIVTM.

Aseptic meningitis syndrome (AMS) may occur with IGIV treatments, including ASCENIVTM. AMS may occur more frequently in association with high doses (2 g/kg) and/or rapid infusion of IGIV.

IGIV products, including ASCENIVTM, may contain blood group antibodies that can act as hemolysins and induce in vivo coating of red blood cells (RBCs) with immunoglobulin, causing a positive direct antiglobulin reaction and hemolysis.

Monitor patients for pulmonary adverse reactions. If TRALI is suspected, perform appropriate tests for the presence of anti-neutrophil antibodies in both the product and the patient's serum.

Because ASCENIVTM is made from human blood, it may carry a risk of transmitting infectious agents, e.g., viruses, the variant Creutzfeldt-Jakob disease (vCJD) and theoretically, the Creutzfeldt-Jakob disease (CJD) agent.

Periodic monitoring of renal function and urine output is particularly important in patients at increased risk of developing acute renal failure. Assess renal function, including measurement of blood urea nitrogen (BUN) and serum creatinine, before the initial infusion of ASCENIVTM and at appropriate intervals thereafter.

After infusion of immunoglobulin, the transitory rise of the various passively transferred antibodies in the patient's blood may yield positive serological testing results, with the potential for misleading interpretation. Passive transmission of antibodies to erythrocyte antigens (e.g., A, B, and D) may cause a positive direct or indirect antiglobulin (Coombs') test.

Adverse Reactions

The most common adverse reactions to ASCENIVTM (\geq 5% of study subjects) were headache, sinusitis, diarrhea, gastroenteritis viral, nasopharyngitis, upper respiratory tract infection, bronchitis, and nausea.

You are encouraged to report side effects of prescription drugs to ADMA Biologics @ 1-800-458-4244 or the FDA. Visit www.fda.gov/MedWatch or call 1-800-FDA-1088.

Cautionary Note Regarding Forward-Looking Statements

This press release contains "forward-looking statements" pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995, about ADMA Biologics, Inc. ("we", "our" or the "Company"). Forward-looking statements include, without limitation, any statement that may predict, forecast, indicate, or imply future results, performance or achievements, and may contain such words as "estimate," "project," "intend," "plan," "planning," "expect," "believe," "will," "is likely," "will likely," "should," "could," "would," "may," or, in each case, their negative, or words or expressions of similar meaning. These forward-looking statements also include, but are not limited to, statements concerning our ability to operate in accordance with FDA quality and compliance, regulations and expectations; our ability to successfully pursue commercialization and prelaunch activities for our products; and the potential of our specialty plasma-based biologics products and product candidates to provide meaningful clinical improvement for patients living with Primary Immune Deficiency Disease or other indications. Actual events or results may differ materially from those described in this document due to a number of important factors. Except to the extent required by applicable laws or rules, ADMA does not undertake any obligation to update any forward-looking statements or to announce revisions to any of the forward-looking statements. Forward-looking statements are subject to many risks, uncertainties and other factors that could cause our actual results, and the timing of certain events, to differ materially from any future results expressed or implied by the forward-looking statements, including, but not limited to, the risks and uncertainties described in our filings with the U.S. Securities and Exchange Commission, including our most recent reports on Form 10-K, 10-Q and 8-K, and any amendments thereto.

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