

Pacira BioSciences Reports Preliminary Second Quarter 2022 Revenue of \$169.4 Million

July 14, 2022

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TAMPA, Fla., July 14, 2022 (GLOBE NEWSWIRE) -- Pacira BioSciences, Inc. (Nasdaq: PCRX), the industry leader in its commitment to non-opioid pain management and regenerative health solutions, today reported preliminary revenues of \$169.4 million for the second quarter of 2022, compared with \$135.6 million for the second quarter of 2021. The company's revenues include net product sales of EXPAREL [®] (bupivacaine liposome injectable suspension), ZILRETTA[®] (triamcinolone acetonide extended-release injectable suspension), and the iovera° system. The company began recognizing sales of ZILRETTA in November 2021 after completing its acquisition of Flexion Therapeutics, Inc.

"We continue to execute our growth strategy and are pleased to have posted record sales in the second quarter. In mid-June, EXPAREL sales trends strengthened despite ongoing pockets of persistent labor shortages and pandemic-related disruptions facing the elective surgery market. This was augmented by ZILRETTA sales, which underscore the successful integration of this synergistic product," said Dave Stack, chairman and chief executive officer of Pacira BioSciences. "EXPAREL has continued to outpace the recovery of the elective surgery market and we are seeing expanding utilization across all target markets and sites of care, with particularly robust growth within outpatient sites of care. We continue to drive education and awareness around the value of ZILRETTA and iovera° as both complementary and standalone non-opioid solutions for managing osteoarthritis pain and remain confident in the long-term growth opportunity for both products."

"On the clinical front, we remain on track with activities advancing according to plan. We have completed enrollment in the first of two studies evaluating EXPAREL as a lower extremity nerve block and we expect to complete enrollment in the second study in the coming weeks. We are also finalizing the design of our label expansion studies for ZILRETTA in shoulder osteoarthritis and type 2 diabetes. In addition, we are preparing to meet with the U.S. Food and Drug Administration (FDA) to discuss the design of our Phase 3 registration study for iovera° as a treatment for spasticity. With a complementary portfolio of innovative non-opioid options, we believe we are uniquely positioned to deliver robust near- and long-term growth," continued Mr. Stack.

Second Quarter and June 2022 Preliminary Revenue Highlights

- EXPAREL net product sales were \$137.0 million and \$130.1 million for the second quarters of 2022 and 2021 and \$47.3 million and \$45.8 million for the months of June 2022 and 2021, respectively. EXPAREL average daily sales were 105 percent of the prior year for the second quarter and 103 percent of the prior year for the month of June, respectively. The company reports average daily growth rates for EXPAREL to account for differences in the number of selling days per reporting period. For the second quarter, EXPAREL selling days were 64 in both 2022 and 2021. For the month of June, EXPAREL selling days were 22 in 2022 and 2021. Sales of bupivacaine liposome injectable suspension to a third-party licensee for use in veterinary practice were \$1.0 million and \$1.0 million in the second quarters of 2022 and 2021, respectively.
- ZILRETTA net product sales were \$27.4 million for the second quarter of 2022 and \$9.8 million for the month of June 2022. ZILRETTA sales in the second quarter of 2021 occurred prior to the completion of the company's acquisition of Flexion in November 2021.
- iovera° net product sales were \$3.2 million and \$3.8 million for the second quarters of 2022 and 2021 and \$1.1 million and \$1.4 million for the months of June 2022 and 2021, respectively.
- Second quarter 2022 royalty revenue was \$0.8 million, compared with \$0.6 million in 2021.

Since early 2020, the company's revenues have been impacted by COVID-19 and pandemic-related challenges that included the significant postponement or suspension in the scheduling of elective surgical procedures due to public health guidance and government directives. While the degree of impact has diminished during the course of the pandemic due to the introduction of vaccines and the lessening of elective surgery restrictions, certain pandemic-related operational challenges persist. It remains unclear how long it will take the elective surgery market to normalize or if restrictions on elective procedures will recur due to future COVID-19 variants or otherwise.

The company is not providing 2022 revenue or gross margin guidance at this time given the continued uncertainty around labor shortages, COVID-19, and the pace of recovery for the elective surgery market. To provide greater transparency, the company is reporting monthly intra-quarter unaudited net product sales for EXPAREL, ZILRETTA, and iovera° until it has gained enough visibility around the impacts of COVID-19. The company is also providing weekly EXPAREL utilization and elective surgery data within its investor presentation, which is accessible at investor.pacira.com. Pacira completed its acquisition of Flexion Therapeutics on November 19, 2021, which added ZILRETTA® (triamcinolone acetonide extended-release injectable suspension) to its commercial offering.

The financial information included in this press release is preliminary, unaudited, and subject to adjustment. It does not present all information necessary for an understanding of the company's financial results for the second quarter or full year 2022.

About Pacira BioSciences

Pacira BioSciences, Inc. (Nasdaq: PCRX) is committed to providing a non-opioid option to as many patients as possible to redefine the role of opioids as rescue therapy only. The company is also developing innovative interventions to address debilitating conditions involving the sympathetic nervous

system, such as cardiac electrical storm, chronic pain, and spasticity. Pacira has three commercial-stage non-opioid treatments: EXPAREL[®] (bupivacaine liposome injectable suspension), a long-acting, local analgesia currently approved for postsurgical pain management; ZILRETTA[®] (triamcinolone acetonide extended-release injectable suspension), an extended-release, intra-articular, injection indicated for the management of osteoarthritis knee pain; and iovera^{o®}, a novel, handheld device for delivering immediate, long-acting, drug-free pain control using precise, controlled doses of cold temperature to a targeted nerve. To learn more about Pacira, including the corporate mission to reduce overreliance on opioids, visit www.pacira.com.

About EXPAREL®

EXPAREL (bupivacaine liposome injectable suspension) is indicated in patients 6 years of age and older for single-dose infiltration to produce postsurgical local analgesia, and in adults as an interscalene brachial plexus nerve block to produce postsurgical regional analgesia. Safety and efficacy have not been established in other nerve blocks. Since its launch, EXPAREL has been used in over nine million patients. EXPAREL utilizes the company's proprietary multivesicular liposomal drug delivery technology composed of a honeycomb of numerous, non-concentric, internal aqueous chambers containing bupivacaine. After injection, bupivacaine is released over time, as the lipid membranes are absorbed, prolonging the duration of action. EXPAREL is the first and only multivesicular liposome local anesthetic that can be utilized in the peri- or postsurgical setting. A single dose of EXPAREL provides significant reductions in cumulative pain scores with up to a 78 percent decrease in opioid consumption; the clinical benefit of the opioid reduction was not demonstrated. Additional information is available at www.EXPAREL.com.

Important Safety Information about EXPAREL for Patients

EXPAREL should not be used in obstetrical paracervical block anesthesia. In studies in adults where EXPAREL was injected into a wound, the most common side effects were nausea, constipation, and vomiting. In studies in adults where EXPAREL was injected near a nerve, the most common side effects were nausea, fever, and constipation. In the study where EXPAREL was given to children, the most common side effects were nausea, vomiting, constipation, low blood pressure, low number of red blood cells, muscle twitching, blurred vision, itching, and rapid heartbeat. EXPAREL can cause a temporary loss of feeling and/or loss of muscle movement. How much and how long the loss of feeling and/or muscle movement depends on where and how much of EXPAREL was injected and may last for up to 5 days. EXPAREL is not recommended to be used in patients younger than 6 years old for injection into the wound, for patients younger than 18 years old for injection near a nerve, and/or in pregnant women. Tell your health care provider if you or your child has liver disease, since this may affect how the active ingredient (bupivacaine) in EXPAREL is eliminated from the body. EXPAREL should not be injected into the spine, joints, or veins. The active ingredient in EXPAREL can affect the nervous system and the cardiovascular system; may cause an allergic reaction; may cause damage if injected into the joints; and can cause a rare blood disorder.

About ZILRETTA®

On October 6, 2017, ZILRETTA (triamcinolone acetonide extended-release injectable suspension) was approved by the U.S. Food and Drug Administration as the first and only extended-release intra-articular therapy for patients confronting osteoarthritis (OA)- related knee pain. ZILRETTA employs proprietary microsphere technology combining triamcinolone acetonide—a commonly administered, short-acting corticosteroid—with a poly lactic-co-glycolic acid (PLGA) matrix to provide extended pain relief. The pivotal Phase 3 trial on which the approval of ZILRETTA was based showed that ZILRETTA significantly reduced OA knee pain for 12 weeks, with some people experiencing pain relief through Week 16. Learn more at www.zilretta.com.

Indication and Select Important Safety Information for ZILRETTA

Indication: ZILRETTA is indicated as an intra-articular injection for the management of OA pain of the knee. Limitation of Use: The efficacy and safety of repeat administration of ZILRETTA have not been demonstrated.

Contraindication: ZILRETTA is contraindicated in patients who are hypersensitive to triamcinolone acetonide, corticosteroids or any components of the product.

Warnings and Precautions:

- Intra-articular Use Only: ZILRETTA has not been evaluated and should not be administered by epidural, intrathecal, intravenous, intraocular, intramuscular, intradermal, or subcutaneous routes. ZILRETTA should not be considered safe for epidural or intrathecal administration.
- Serious Neurologic Adverse Reactions with Epidural and Intrathecal Administration: Serious neurologic events have been reported following epidural or intrathecal corticosteroid administration. Corticosteroids are not approved for this use.
- **Hypersensitivity reactions:** Serious reactions have been reported with triamcinolone acetonide injection. Institute appropriate care if an anaphylactic reaction occurs.
- **Joint infection and damage:** A marked increase in joint pain, joint swelling, restricted motion, fever and malaise may suggest septic arthritis. If this occurs, conduct appropriate evaluation and if confirmed, institute appropriate antimicrobial treatment.

Adverse Reactions: The most commonly reported adverse reactions (incidence ≥1%) in clinical studies included sinusitis, cough, and contusions.

Please see ZILRETTALabel.com for full Prescribing Information.

About iovera®

The iovera° system is used to destroy tissue during surgical procedures by applying freezing cold. It can also be used to produce lesions in peripheral nervous tissue by the application of cold to the selected site for the blocking of pain. It is also indicated for the relief of pain and symptoms associated with osteoarthritis of the knee for up to 90 days. In one study, the majority of the patients suffering from osteoarthritis of the knee experienced pain and system relief beyond 150 days. The iovera° system's "1×90" Smart Tip configuration (indicating one needle which is 90 mm long) can also facilitate target nerve location by conducting electrical nerve stimulation from a separate nerve stimulator. The iovera° system is not indicated for treatment of

central nervous system tissue. Additional information is available at www.iovera.com.

Important Safety Information for iovera®

The iovera° system is contraindicated for use in patients with the following: Cryoglobulinemia; Paroxysmal cold hemoglobinuria; cold urticaria; Raynaud's disease; open and/or infected wounds at or near the treatment line. Potential complications: As with any surgical treatment that uses needle-based therapy, there is potential for temporary site-specific reactions, including but not limited to: bruising (ecchymosis); swelling (edema); inflammation and/or redness (erythema); pain and/or tenderness; altered sensation (localized dysesthesia). Typically, these reactions resolve with no physician intervention. Patients may help the healing process by applying ice packs to the affected sites, and by taking over-the-counter analgesics.

Forward-Looking Statements

Any statements in this press release about Pacira's future expectations, plans, trends, outlook, projections and prospects, and other statements containing the words "believes," "anticipates," "plans," "estimates," "expects," "intends," "may," "will," "would," "could," "could," "can" and similar expressions, constitute forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. These forward-looking statements include, without limitation, statements related to the acquisition of Flexion Therapeutics, Inc. and the costs and benefits thereof, our growth and future operating results and trends, our strategy, plans, objectives, expectations (financial or otherwise) and intentions, future financial results and growth potential, anticipated product portfolio, development programs, strategic alliances, patent terms and intellectual property and other statements that are not historical facts. For this purpose, any statement that is not a statement of historical fact should be considered a forward-looking statement. We cannot assure you that our estimates, assumptions and expectations will prove to have been correct. Actual results may differ materially from these indicated by such forward-looking statements as a result of various important factors, including risks relating to, among others: risks associated with acquisitions, such as the risk that the businesses will not be integrated successfully, that such integration may be more difficult, time-consuming or costly than expected or that the expected benefits of the transaction will not occur; the possibility that if we do not achieve the perceived benefits of the Flexion acquisition as rapidly or to the extent anticipated by financial analysts or investors, the market price of our shares could decline; the impact of the COVID-19 pandemic on elective surgeries, our manufacturing and supply chain, global and United States economic conditions, and our business, including our revenues, financial condition and results of operations; the success of our sales and manufacturing efforts in support of the commercialization of EXPAREL, ZILRETTA and iovera° and the rate and degree of market acceptance of EXPAREL, ZILRETTA and iovera°; the size and growth of the potential markets for EXPAREL, ZILRETTA and iovera° and our ability to serve those markets; our plans to expand the use of EXPAREL, ZILRETTA and iovera° to additional indications and opportunities, and the timing and success of any related clinical trials for EXPAREL, ZILRETTA and iovera°; the commercial success of EXPAREL, ZILRETTA and iovera°; the related timing and success of United States Food and Drug Administration supplemental New Drug Applications, and premarket notification 510(k)s; the related timing and success of European Medicines Agency Marketing Authorization Applications; our plans to evaluate, develop and pursue additional product candidates utilizing our proprietary multivesicular liposome (pMVL) drug delivery technology; the approval of the commercialization of our products in other jurisdictions; clinical trials in support of an existing or potential pMVL-based product; our commercialization and marketing capabilities, our ability to successfully construct an additional EXPAREL manufacturing suite in San Diego, California; our ability to successfully complete a ZILRETTA capacity expansion project in Swindon, England; the outcome of any litigation; the ability to successfully integrate Flexion or any future acquisitions into our existing business; the recoverability of our deferred tax assets; and assumptions associated with contingent consideration payments; and factors discussed in the "Risk Factors" of our most recent Annual Report on Form 10-K and in other filings that we periodically make with the Securities and Exchange Commission. In addition, the forward-looking statements included in this press release represent our views as of the date of this press release. Important factors could cause actual results to differ materially from those indicated or implied by forward-looking statements, and as such we anticipate that subsequent events and developments will cause our views to change. However, while we may elect to update these forward-looking statements at some point in the future, we specifically disclaim any obligation to do so, except as required by law. These forward-looking statements should not be relied upon as representing our views as of any date subsequent to the date of this press release.

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