## Clarivate

## Tips for tracking the impact of COVID-19 on clinical research and help you navigate the uncertainties ahead

## **Cortellis Clinical Trials Intelligence**

Track clinical trials that are suspended or with delayed patient enrollment due to COVID-19

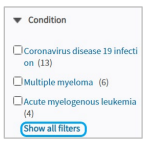
1. Type Covid-19 AND (delay OR halt OR pause OR postpone) in the Quick Search box and click the Full Text search button

```
All Covid-19 AND (delay OR halt OR pause OR postpone)
```

Index Q

Full Text Q

2. Under Report Type (on the left panel), switch to **Clinical Trials**, then click the **Show all Filters** link below Condition.



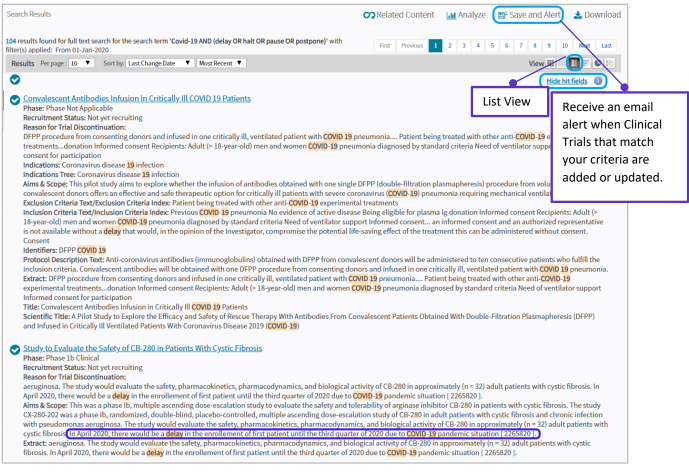
3. Scroll to the last filter to define the **Last Change Date** from January 1, 2020 and leave the to blank. Apply the filters.

Optional Filters: You may also select clinical trials in your **Conditions** of interest or those from specific **Sponsors**. If you switch the **Recruitment Status** filer, you will see which clinical studies officially got Suspended or Terminated, as well as Planned studies that are now delayed due to COVID-19.

4. On the Search Results\* page, click on the **List View** thumbnail (on the top right) then **Show hit fields**\* to view where the search terms appear in each report. You may also **Sort by** Last Change Date of Most Recent.

Use the **Save and Alert** feature to receive email notifications on any clinical trials that get disrupted, as well as any adjustments to the protocol and estimated end dates.





1. Explore the results.

This will include Suspended or Terminated clinical trials and even Planned studies that have been delayed, as well as treatments in development for Coronavirus disease 19 infection.

 To view comments regarding the type of impact on these trials, under Protocols & Results, go to the Aims & Scope section. You may also search (Ctrl +F) for where COVID-19 is mentioned.

## Clarivate<sup>®</sup>

< Back   Forward > C	Clinical Trial Report					🗘 Alert	📥 Dow	
Study to Evaluate the Safety	of CB-280 in Patients With Cystic Fibrosi	S			Prev	Next 🕨		
Snapshot	Highlight 🔲 Search Terms & Synonyms	< Previous	Next >		PRESS RELEASE		×	
Protocol & Results	PROTOCOL & RESULTS				14-Apr-2020			
Aims & Scope	AIMS & SCOPE This was a phase lb, multiple ascending dose-escalation study to evaluate the safety and tolerability of arginase inhibitor CB-280 in P Calibhera Biosciences Provides Update on Business Operations Calibhera Biosciences Inc							
Protocol Description     Regimens	The study CX-280-202 was a phase lb, randomized, double-blind, placebo-controlled, multiple ascending dose-escalation study of CE chronic infection with pseudomonas aeruginosa. The study would evaluate the safety, pharmacokinetics, pharmacokynamics, and b							
Results	In April 2020, there would be a delay in the enrollement of first patient until the third quarter of 2020 due to COVID-19 pandemic situation [2265820].							
<ul> <li>Adverse Events</li> </ul>	PROTOCOL DESCRIPTION TEXT							
Treatment	There would be four planned sequential dose-escalation, cohorts of eight patients each, randomized 6 : 2 to receive CB-280 or matched placebo at doses of 50, 100, 200, or 400 mg po, administered bid for 14 days and placebo twice-daily for 14 days. Intermediate dose levels might be evaluated based on emerging safety data at the planned dose levels.							
Subjects & Measurements			-			^		
Registry Contacts & Sites								
Change History								
Sources								

3. Click on the blue reference numbers to see the source. For Press Releases, you may review the full report in Cortellis to get more details such as what measures the company is taking, any new data read out dates, etc.

Calithera Biosciences Provides Update on Business Operations									
Snapshot	Highlight 🔲 Search Terms & Synonyms	< Previous	Next >						
Full Text	FULL TEXT								
	April 14, 2020 07:00 ET   Source: Calithera Biosciences, Inc.								
	SOUTH SAN FRANCISCO, April 14, 2020 (GLOBE NEWSWIRE) – Calithera Biosciences, Inc. (Nasdaq: CALA), a clinical stage biotechnology company focused on discovering and developing novel small-molecule drugs for the treatment of cancer and other life-threatening diseases, shared an update on business operations, including clinical program adjustments related to COVID-19.								
	"Our first priority during the COVID-19 pandemic is the health of our employees, as well as patients and medical professionals involved in our clinical programs. We are continuing all clinical operations, with additional COVID-19 related safety measures in place. Given the fluid nature of the current situation, and the impact of the pandemic on clinical sites globally, we are delaying the start of enrollment of patients in our two new clinical trials, while we work towards accelerating the opening of sites," said Susan Molineaux, president and chief executive officer of Calithera. "We believe we are on track to announce top-line CANTATA results this year, and we are narrowing the timeframe to the fourth quarter of 2020."								
	Clinical programs update:								
	Randomized CANTATA trial of telaglenastat and cabozantinib in advanced renal cell carcinoma. The CANTATA trial was fully enrolled in October 2019 and Calithera advised at that time that the company planned to report top-line at fastev data from the trial in the second half of 2020, and more recently guided towards late third quarter of routh quarter of 2020. In light of COMD-19, Calithera now expects top-line data in the fourth quarter of 2020. Calithera has made accommodations to facilitate study conduct during the pandemic, including allowing patients to have scans performed at local clinical centers to facilitate compliance with the study schedule of assessments, and to receive a larger allocation of study drug in order to reduce the number of visits required to the clinical site, if necessary. While affirming that the readout is expected by the end of 2020, the updated guidance allows for additional time for activities that required to the clinical site, if necessary. While affirming that the readout is expected by the end of 2020, the updated guidance allows for additional time for activities that required to the clinical site in fourthing data monitoring.								
	lung cancer continues to progress towards mult	KE trial in non-small cell lung cancer (NSCLC) patients with genetic mutation NRF2/KEAP1. The randomized Phase 2 trial of telaglenastat for the treatment of to progress towards multiple site openings. However, given the challenges associated with opening new clinical studies during the current stage of the COVID- a expects to delay enrollment of the first patient until the third quarter of 2020, pending further developments in the COVID-19 situation. Calithera plans to rom this trial in 2021.							

For more information or to request for a guided workflow review, contact Customer Service at LS Product Support.