

Patent Office Reduces Accelerated Examination Fees for COVID-19 Treatments

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(by [Carl Ronald](#))

Not only has the global pandemic spawned a race to develop a cure for COVID-19, it has also created a race to the Patent Office to protect the massive investments companies are making in their attempts to develop novel diagnostics, therapies, and vaccines to combat the disease. In the United States, the first inventor to file a patent application that teaches a new and non-obvious way to treat the virus or its effects will be eligible for the limited monopoly that a patent provides. As has been reported, researchers are employing a variety of different mechanisms to attack the virus and it is expected that a significant number of patents will ultimately issue from these efforts.



While Big Pharma is well-poised to incur the expense of patent filing, including the additional expense of paying for accelerated examination of their applications, smaller concerns may not be in a position to do so. To help small businesses and solo inventors in this regard, the United States Patent Office has announced a [program](#) that would fast-track the examination of certain patent applications related to the pandemic. While the typical turn-around time for an Examiner to provide an initial review of a new application is about two years after filing, payment of an extra fee to accelerate examination is also an option. Small businesses and

startup companies, however, typically can't take advantage of this program due to the substantial increased initial cost. To remove this impediment, the new program enables business with less than 500 employees to request accelerated examination of certain COVID-19-related applications with no additional upfront payment. If an application qualifies for the program, the Patent Office promises to fully examine it within a year of being granted prioritized status.

The program is further limited to therapeutics, vaccines and diagnostic tests that are subject to FDA approval for COVID-19 related uses. FDA approvals include, for example, Investigational New Drug (IND) applications, Investigational Device Exemptions (IDE), New Drug Applications (NDA), Biologics License Applications (BLA), Pre-market Approvals (PMA), and Emergency Use Authorizations (EUA).

As of Thursday, August 20th, there have been 274 applicants with 146 requests having been granted. The program is limited to the first 500 approved applicants.

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