

# PMPRB's decision that patent pertains to drug held to be reasonable by Federal Court

January 23, 2024

In [Galderma Canada Inc. v. Canada \(Attorney General\)](#), the Federal Court (FC) judicially reviewed a decision relating to a drug that contains 0.1 per cent of a medicine while the patent is to a 0.3 per cent concentration of that medicine.

Galderma and the Patented Medicine Prices Review Board (PMPRB) have a long-running dispute over whether a particular patent “pertains to” Galderma’s Differin® product, under the Patent Act. In the most recent [judicial review decision](#), the FC held that the PMPRB’s decision that the patent does pertain to the product was reasonable.

## History

Galderma sells two products containing the medicinal ingredient adapalene: Differin® contains 0.1 per cent adapalene whereas Differin XP® contains 0.3 per cent. The patents pertaining to the Differin® product had expired, meaning its pricing was no longer under the PMPRB’s jurisdiction. The ‘237 Patent, which is the subject of this dispute, has also now expired. However, the dispute is about whether it pertained to the Differin® product prior to expiry, bringing that product back under PMPRB jurisdiction while the patent was in force.

In 2016, the [PMPRB determined that the ‘237 Patent does pertain to Differin®](#). However, in 2017, the [FC quashed that decision](#), holding that it was unreasonable for the Board, without explanation, to conclude that a patent relating to a composition of 0.3 per cent adapalene can be used for a medicine with a composition of only 0.1 per cent adapalene.

The [Attorney General’s appeal to the Federal Court of Appeal \(FCA\) was granted](#), remitting the decision to the Board for redetermination. The FCA held that the invention of the ‘237 Patent is a composition with a 0.3 per cent concentration of adapalene for the treatment of dermatological disorders.

Furthermore, the FCA held that the metaphor of the “merest slender thread” cannot replace the statutory definition of “pertains to” in the Patent Act. The FCA held that “[i]n cases such as this, where the question is whether an invention pertains to a specific

medicine, what kind of clinical similarities would support a finding that the invention of a patent was intended or capable of being used for that medicine?” (para 73).

The parties submitted further written representations before the Board, but no new evidence. In May 2020, [the PMPRB again held that the ‘237 Patent pertains to Differin®](#). The Board held that Galderma’s argument that for an invention to pertain to a medicine under the Patent Act, it must encompass the medicine being sold by the patentee, was inconsistent with the wording of the provision and precluded by the decision of the FCA. Furthermore, due to the clinical similarities between the two products, the Board was satisfied that the invention in the ‘237 Patent could be used for Differin®.

As part of the judicial review, Galderma filed three additional affidavits as evidence before the FC, those of an expert in patent law, a regulatory affairs expert, and a fact witness. On motion, the [Court struck the affidavit of the patent expert in full, and the regulatory affairs expert's in part](#). The fact witness' affidavit was allowed as it contained non-controversial background information.

## The Federal Court decision

The FC concluded that the standard of review is reasonableness, which is the same standard previously applied by the FCA. [Citing the Supreme Court](#), the FC held it is only to intervene where “there are sufficiently serious shortcomings in the decision such that it cannot be said to exhibit the requisite degree of justification, intelligibility and transparency.” Thus, the reasons must allow an understanding of why the decision was made, as well as a determination of whether it falls within the range of acceptable outcomes defensible in respect of the facts and law (para 34).

The FC dismissed Galderma’s argument that the Board’s decision was not procedurally fair, as the product monograph (PM) was not mentioned in the Board’s Notice of Application. Galderma did not object to this alleged breach before the Board. Furthermore, the PM had been referred to in the Board’s initial decision, the FC’s previous decision, and by the FCA.

The FC held that the Board’s decision that Differin® and Differin XP® “use the same medicinal ingredient, are indicated for the same dermatological disorder and work in the same way,” and thus are the same medicine, was reasonably supported by the evidence (para 60). The FC also took no issue with the Board’s conclusions that the shared PM supported the existence of a rational connection between the two products. Furthermore, the invention of the ‘237 Patent and Differin® produced similar clinical effects, with comparable side effects, and while not interchangeable, they were prescribed for similar conditions and could in some circumstances be substituted for each other.

The FC concluded that the FCA had remitted a narrow issue for redetermination and the PMPRB needed to “consider the kind of clinical similarities that would support a finding that the invention of a patent was intended or capable of being used for that medicine” (para 64). The Board found significant clinical similarities, and its decision was reasonable.

Thus, the judicial review was dismissed, with lump-sum costs payable to the Attorney General of Canada.

By

Beverley Moore, Chantal Saunders

Expertise

Disputes, Intellectual Property, Health Care & Life Sciences

---

## **BLG | Canada's Law Firm**

As the largest, truly full-service Canadian law firm, Borden Ladner Gervais LLP (BLG) delivers practical legal advice for domestic and international clients across more practices and industries than any Canadian firm. With over 800 lawyers, intellectual property agents and other professionals, BLG serves the legal needs of businesses and institutions across Canada and beyond – from M&A and capital markets, to disputes, financing, and trademark & patent registration.

[blg.com](http://blg.com)

## **BLG Offices**

### **Calgary**

Centennial Place, East Tower  
520 3rd Avenue S.W.  
Calgary, AB, Canada  
T2P 0R3

T 403.232.9500  
F 403.266.1395

### **Ottawa**

World Exchange Plaza  
100 Queen Street  
Ottawa, ON, Canada  
K1P 1J9

T 613.237.5160  
F 613.230.8842

### **Vancouver**

1200 Waterfront Centre  
200 Burrard Street  
Vancouver, BC, Canada  
V7X 1T2

T 604.687.5744  
F 604.687.1415

### **Montréal**

1000 De La Gauchetière Street West  
Suite 900  
Montréal, QC, Canada  
H3B 5H4

T 514.954.2555  
F 514.879.9015

### **Toronto**

Bay Adelaide Centre, East Tower  
22 Adelaide Street West  
Toronto, ON, Canada  
M5H 4E3

T 416.367.6000  
F 416.367.6749

The information contained herein is of a general nature and is not intended to constitute legal advice, a complete statement of the law, or an opinion on any subject. No one should act upon it or refrain from acting without a thorough examination of the law after the facts of a specific situation are considered. You are urged to consult your legal adviser in cases of specific questions or concerns. BLG does not warrant or guarantee the accuracy, currency or completeness of this publication. No part of this publication may be reproduced without prior written permission of Borden Ladner Gervais LLP. If this publication was sent to you by BLG and you do not wish to receive further publications from BLG, you may ask to remove your contact information from our mailing lists by emailing [unsubscribe@blg.com](mailto:unsubscribe@blg.com) or manage your subscription preferences at [blg.com/MyPreferences](http://blg.com/MyPreferences). If you feel you have received this message in error please contact [communications@blg.com](mailto:communications@blg.com). BLG's privacy policy for publications may be found at [blg.com/en/privacy](http://blg.com/en/privacy).

© 2026 Borden Ladner Gervais LLP. Borden Ladner Gervais LLP is an Ontario Limited Liability Partnership.