

FORM 52-109F2R
CERTIFICATION OF REFILED INTERIM FILINGS

This certificate is being filed on the same date that ***Titan Medical Inc.***, “issuer”) has refiled ***The Management and Discussion Analysis***.

I, ***David McNally, Chief Executive Officer***, certify the following:

1. ***Review:*** I have reviewed the interim financial statements and interim MD&A (together, the “interim filings”) of the issuer for the interim period ended ***September 30, 2018***.
2. ***No misrepresentations:*** Based on my knowledge, having exercised reasonable diligence, the interim filings do not contain any untrue statement of a material fact or omit to state a material fact required to be stated or that is necessary to make a statement not misleading in light of the circumstances under which it was made, with respect to the period covered by the interim filings.
3. ***Fair presentation:*** Based on my knowledge, having exercised reasonable diligence, the interim financial report together with the other financial information included in the interim filings fairly present in all material respects the financial condition, financial performance and cash flows of the issuer, as of the date of and for the periods presented in the interim filings.
4. ***Responsibility:*** The issuer’s other certifying officer and I are responsible for establishing and maintaining disclosure controls and procedures (DC&P) and internal control over financial reporting (ICFR), as those terms are defined in National Instrument 52-109 *Certification of Disclosure in Issuers’ Annual and Interim Filings*, for the issuer.
5. ***Design:*** Subject to the limitations, if any, described in paragraphs 5.2 and 5.3, the issuer’s other certifying officer and I have, as at the end of the period covered by the interim filings
 - (a) designed DC&P, or caused it to be designed under our supervision, to provide reasonable assurance that
 - (i) material information relating to the issuer is made known to us by others, particularly during the period in which the interim filings are being prepared; and
 - (ii) information required to be disclosed by the issuer in its annual filings, interim

filings or other reports filed or submitted by it under securities legislation is recorded, processed, summarized and reported within the time periods specified in securities legislation; and

- (b) designed ICFR, or caused it to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with the issuer's GAAP.

- 5.1 **Control framework:** The control framework the issuer's other certifying officer and I used to design the issuer's ICFR is Integrated Framework (COSO).
- 5.2 **ICFR – material weakness relating to design:** N/A
- 5.3 **Limitation on scope of design:** N/A
- 6. **Reporting changes in ICFR:** The issuer has disclosed in its interim MD&A any change in the issuer's ICFR that occurred during the period beginning on January 1, 2018 and ended on September 30, 2018 that has materially affected, or is reasonably likely to materially affect, the issuer's ICFR.

Date: November 12, 2018

(signed) "David McNally"

David McNally
Chief Executive Officer
Titan Medical Inc.

Date: **November 16, 2018**

(signed) "David McNally"

Chief Executive Officer