PATENT

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Applicant(s) Myers et al. Examiner: Janet L. Epps-Smith

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For: Sublingual and Buccal Film

Compositions

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Dated: February 29, 2012

Signature: Christine Briscoe/cbriscoe/

AMENDMENT AND RESPONSE

Sir:

In response to the office action dated August 31, 2011, a response to which is due by February 29, 2012 in view of the concurrently filed petition for three month extension of time, please amend the application as follows:

Amendments to the Claims begin on page 2 of this paper.

Remarks begin on page 7 of this paper.



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Amendments to the Claims:

This listing of claims shall replace all previous listings in this application:

- 1. (Currently Amended) A film dosage composition comprising:
 - a. A polymeric carrier matrix;
 - b. A therapeutically effective amount of buprenorphine or a pharmaceutically acceptable salt thereof;
 - c. A therapeutically effective amount of naloxone or a pharmaceutically acceptable salt thereof; and
 - d. A buffer in an amount to provide a local pH of for said composition of a value sufficient to optimize absorption of said buprenorphine, wherein said local pH is from about 2 to about 3.5 in the presence of saliva.
- 2. (Canceled).
- 3. (Currently Amended) The composition of claim <u>1</u> 2, wherein the local pH of said composition is from about 3 to about <u>3.5</u> [[4]].
- 4. (Original) The composition of claim 1, wherein said film dosage composition provides a bioequivalent absorption of buprenorphine to that of a tablet having an equivalent amount of buprenorphine or a pharmaceutically acceptable salt thereof.
- 5. (Original) The composition of claim 1, wherein said polymeric carrier matrix comprises at least one polymer in an amount of at least 25% by weight of said composition.
- 6. (Original) The composition of claim 1, wherein said buffer is present in an amount of from about 2:1 to about 1:5 by weight of buffer to buprenorphine.
- 7. (Original) The composition of claim 1, wherein said polymeric carrier matrix comprises at least one self-supporting film forming polymer.
- 8. (Original) The film dosage composition of claim 1, wherein said buprenorphine is present in an amount of from about 2 mg to about 16 mg per dosage.
- 9. (Original) The film dosage composition of claim 1, wherein said buffer comprises sodium citrate, citric acid, and combinations thereof.



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- 10. (Original) The film dosage composition of claim 1, wherein said buffer comprises acetic acid, sodium acetate, and combinations thereof.
- 11. (Currently Amended) A film dosage composition comprising:
 - a. A polymeric carrier matrix;
 - b. A therapeutically effective amount of buprenorphine or a pharmaceutically acceptable salt thereof;
 - c. A therapeutically effective amount of naloxone or a pharmaceutically acceptable salt thereof; and
 - d. A buffer in an amount sufficient to inhibit the absorption of said naloxone, while also optimizing absorption of said buprenorphine when administered orally.
- 12. (Currently Amended) The composition of claim 11, wherein said composition has a local pH of about 2 to about 3.5 [[4]].
- 13. (Currently Amended) The composition of claim 11, wherein <u>said composition has a local pH of about 3 to about 3.5 said buffer is present in an amount sufficient to provide a therapeutically adequate absorption of buprenorphine.</u>
- 14. (Currently Amended) The composition of claim 13, wherein a therapeutically adequate absorption of buprenorphine comprises said buffer is present in an amount sufficient to provide-a bioequivalent level of absorption of buprenorphine as a tablet having an equivalent amount of buprenorphine or a pharmaceutically acceptable salt thereof.
- 15. (Currently Amended) A film dosage composition comprising:
 - a. A polymeric carrier matrix;
 - b. A therapeutically effective amount of buprenorphine or a pharmaceutically acceptable salt thereof;
 - c. A therapeutically effective amount of naloxone or a pharmaceutically acceptable salt thereof; and
 - d. A buffering system;

wherein said buffering system comprises a buffer capacity sufficient to maintain the ionization of naloxone during the time which said composition is in the oral



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cavity of a user, and also sufficient to optimize the absorption of said buprenorphine.

- 16. (Currently Amended) The composition of claim 15, wherein said composition has a local pH of about 2 to about 3.5 [[4]].
- 17. (Currently Amended) A method of treating narcotic dependence of a user, comprising the steps of:
 - a. providing a composition comprising:
 - i. A polymeric carrier matrix;
 - ii. A therapeutically effective amount of buprenorphine or a pharmaceutically acceptable salt thereof;
 - iii. A therapeutically effective amount of naloxone or a pharmaceutically acceptable salt thereof; and
 - iv. A buffer in an amount to provide a local pH of <u>about 2 to about 3.5 for</u> said composition of a value sufficient to optimize absorption of said buprenorphine <u>and also sufficient to inhibit absorption of said naloxone</u>; and
 - b. administering said composition to the oral cavity of a user.
- 18. (Original) The composition of claim 17, wherein said method provides a bioequivalent absorption of buprenorphine to that of a tablet having an equivalent amount of buprenorphine or a pharmaceutically acceptable salt thereof.
- 19. (Currently Amended) The method of claim 17, wherein said composition has a local pH of about 3 [[2]] to about 3.5 [[4]].
- 20. (Original) The method of claim 17, wherein said film dosage composition is administered to the user through buccal administration, sublingual administration, and combinations thereof.
- 21. (Original) The method of claim 17, wherein said film dosage composition remains in the oral cavity of the user for a period of at least 1 minute.
- 22. (Original) The method of claim 17, wherein said film dosage composition remains in the oral cavity of the user for a period of between about 1 and 1.5 minutes.



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- 23. (Original) The method of claim 17, wherein said film dosage composition remains in the oral cavity of the user for a period of up to 3 minutes.
- 24. (Currently Amended) A process of forming a film dosage composition comprising the steps of:
 - a. casting a film-forming composition, said film-forming composition comprising:
 - i. A polymeric carrier matrix;
 - ii. A therapeutically effective amount of buprenorphine or a pharmaceutically acceptable salt thereof;
 - iii. A therapeutically effective amount of naloxone or a pharmaceutically acceptable salt thereof; and
 - iv. A buffer in an amount to provide a local pH of said composition of a
 value sufficient to optimize absorption of said buprenorphine and also
 sufficient to inhibit absorption of said naloxone; and
 - b. drying said film-forming composition to form a self-supporting film dosage composition.
- 25. (Currently Amended) The process of claim 24, wherein said composition has a local pH of about 2 to about 3.5 [[4]].
- 26. (Currently Amended) A film dosage composition comprising a therapeutically sufficient amount of buprenorphine or a pharmaceutically acceptable salt thereof and a therapeutically sufficient amount of naloxone or a pharmaceutically acceptable salt thereof, said film dosage composition having a bioequivalent release profile as a tablet containing about 2 times the amount of buprenorphine or a pharmaceutically acceptable salt thereof, and wherein said composition provides a local pH of from about 2 to about 3.5.
- 27. (Original) An orally dissolving film formulation comprising buprenorphine and naloxone, wherein said formulation provides an in vivo plasma profile having a Cmax of between about 0.624 ng/ml and about 5.638 ng/ml for buprenorphine and an in vivo plasma profile having a Cmax of between about 41.04 pg/ml to about 323.75 pg/ml for naloxone.



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