

STATE OF NORTH CAROLINA
WAKE COUNTY

**FILED IN THE GENERAL COURT OF JUSTICE
SUPERIOR COURT DIVISION**

2014 JUN -4 FILE NO. CVS - 7276

WAKE COUNTY, C.S.C.

STATE OF NORTH CAROLINA, *ex rel.*)
ROY COOPER, ATTORNEY GENERAL,)

Plaintiff,)

vs.)

GLAXOSMITHKLINE LLC,)

Defendant.)

CONSENT JUDGMENT

THIS CAUSE came on before the undersigned Superior Court Judge for entry of a Consent Judgment between Plaintiff, State of North Carolina, by and through its Attorney General, and Defendant GlaxoSmithKline LLC (hereinafter "GlaxoSmithKline" or "GSK"). The Court finds that these two parties have resolved the matters in controversy between them and have agreed to the entry of this Consent Judgment ("Consent Judgment") by the Court without trial or adjudication of any issue of fact or law, and without finding or admission of wrongdoing or liability of any kind.

IT IS HEREBY ORDERED THAT:

I. FINDINGS AND CONCLUSIONS

FINDINGS OF FACT

A. The State of North Carolina, by and through its Attorney General, is the Plaintiff in this case. The North Carolina Attorney General is charged with, among other things, the

responsibility of enforcing the Unfair and Deceptive Trade Practices Act, N.C. Gen. Stat. § 75-1.1 *et seq.*

B. Plaintiff has filed a Complaint for a permanent injunction and other relief in this matter pursuant to N.C. Gen. Stat. § 75-1.1, alleging that GlaxoSmithKline committed violations of the aforementioned Act.

C. GlaxoSmithKline, at all times relevant hereto, engaged in trade and commerce affecting consumers within the meaning of N.C. Gen. Stat. § 75-1.1 *et seq.* in the State of North Carolina, including but not limited to Wake County.

D. The Attorneys General conducted an investigation regarding the Covered Conduct. The Parties have agreed to resolve all issues raised by and concerns related to the Covered Conduct under N.C. Gen. Stat. § 75-1.1 by entering into this Consent Judgment.

CONCLUSIONS OF LAW

A. This Court has jurisdiction over the subject matter of this lawsuit and over all Parties.

B. The terms of this Consent Judgment shall be governed by the laws of the State of North Carolina.

C. Good cause exists for the Court to enter judgment as to Defendant.

D. Entry of this Consent Judgment is in the public interest and reflects a negotiated agreement between the Parties.

E. Defendant has, by signature of its counsel hereto, waived any right to appeal, petition for certiorari, or move to reargue or rehear this Consent Judgment.

F. This Consent Judgment is just and reasonable with respect to all Parties. The Court approves the terms of the Parties' agreement and adopts them as its own determination of the Parties' respective rights and obligations.

G. The Attorneys General conducted an investigation regarding the Covered Conduct. The Parties have agreed to resolve the concerns related to the Covered Conduct under the relevant State Consumer Protection Laws by entering into this Consent Judgment. This Consent Judgment reflects a negotiated agreement entered into by the Parties as their own free and voluntary act, and with full knowledge and understanding of the nature of the proceedings and the obligations and duties imposed by this Consent Judgment. GSK is entering into this Consent Judgment solely for the purpose of settlement, and nothing contained herein may be taken as or construed to be an admission or concession of any violation of law or regulation, or of any other matter of fact or law, or of any liability or wrongdoing, all of which GSK expressly denies. Through this Consent Judgment, GSK does not admit any violation of law, and does not admit any wrongdoing that was or could have been alleged by any of the signatory Attorneys General before the date of the Consent Judgment. No part of this Consent Judgment, including its statements and commitments, shall constitute evidence of any liability, fault, or wrongdoing by GSK. This Consent Judgment does not constitute an admission by GSK that the Covered Conduct violated or could violate the Relevant State Consumer Protection Laws. It is the intent of the Parties that this Consent Judgment shall not be admissible or binding in any other matter, including, but not limited to, any investigation or litigation, other than in connection with the enforcement of this Consent Judgment. No part of this Consent Judgment shall create a private cause of action or convert any right to any third party for violation of any federal or state statute or law, except that an Attorney General may file an action to enforce the terms of this Consent

Judgment. Nothing contained herein prevents or prohibits the use of this Consent Judgment for purposes of enforcement by the North Carolina Attorney General.

H. This Consent Judgment does not create a waiver or limit GSK's legal rights, remedies, or defenses in any other action by the North Carolina Attorney General, and does not waive or limit GSK's right to defend itself from, or make arguments in, any other matter, claim, or suit, including, but not limited to, any investigation or litigation relating to the existence, subject matter, or terms of this Consent Judgment. Nothing in this Consent Judgment shall waive, release, or otherwise affect any claims, defenses, or other positions GSK may assert in connection with any investigations, claims, or other matters the Attorneys General are not releasing hereunder. Notwithstanding the foregoing, the North Carolina Attorney General may file an action to enforce the terms of this Consent Judgment.

I. This Consent Judgment does not constitute an approval by the Attorneys General of GSK's business practices, and GSK shall make no representation or claim to the contrary.

J. This Consent Judgment sets forth the entire agreement between the Parties hereto and supersedes all prior agreements or understandings, whether written or oral, between the Parties and/or their respective counsel, with respect to the Covered Conduct.

K. This Court retains jurisdiction of this Consent Judgment and the Parties hereto for the purpose of enforcing and modifying this Consent Judgment and for the purpose of granting such additional relief as may be necessary and appropriate.

L. This Consent Judgment may be executed in counterparts, each of which shall be deemed to constitute an original counterpart hereof, and all of which shall together constitute one and the same Consent Judgment. One or more counterparts of this Consent Judgment may be delivered by facsimile or electronic transmission with the intent that it, or they, shall constitute

an original counterpart hereof. This Consent Judgment relates solely to GSK's business in the United States.

M. This Consent Judgment (or any portion thereof) shall in no way be construed to prohibit GSK from making representations with respect to any GSK Product that are permitted under Federal law or labeling for the drug under the most current draft or final standard promulgated by the FDA or the most current draft or final FDA Guidance for Industry, or permitted or required under any IND, NDA, sNDA, or ANDA approved by the FDA, so long as the representation, taken in its entirety, is not false, misleading or deceptive.

N. Nothing in this Judgment shall require GSK to:

- a) take any action that is prohibited by the FDCA or any regulation promulgated thereunder, or by the FDA; or
- b) fail to take any action that is required by the FDCA or any regulation promulgated thereunder, or by the FDA;

or shall preclude GSK from providing health care economic information to a formulary committee or similar entity or its members in the course of the committee or entity carrying out its responsibilities for the selection of drugs for managed care or other similar organizations pursuant to the standards of Section 114 of the Food and Drug Administration Modernization Act of 1997 (FDAMA), if the information directly relates to an approved indication of a GSK Product, and if based on competent and reliable scientific evidence.

II. DEFINITIONS

The following definitions shall be used in construing this Consent Judgment:

1. "Applicable Clinical Trials" shall mean those clinical trials required by the FDA Amendments Act of 2007 (Public Law No. 110-85).

2. "Attorneys General" shall mean the Attorneys General of the Multistate Working Group.
3. "Clinically Relevant Information" shall mean information that reasonably prudent clinicians would consider relevant when making prescribing decisions regarding a GSK Product.
4. "Clinical Response" shall mean a non-Promotional, scientific communication to address Unsolicited Requests for medical information.
5. "Covered Conduct" shall mean GSK's Promotional practices, dissemination of information, and remuneration to HCPs regarding the prescription drugs Advair®, Paxil®, and Wellbutrin® in the United States.
6. "Effective Date" shall mean the date on which a copy of this Consent Judgment, duly executed by GSK and by the signatory Attorney General, is approved by, and becomes a Judgment, of the Court.
7. "GlaxoSmithKline LLC," "GlaxoSmithKline," or "GSK" shall mean GlaxoSmithKline LLC, including all of its predecessors, subsidiaries, successors, and assigns.
8. "GSK Law Department" shall mean personnel of the GSK Law Department or its designee providing legal advice to GSK.
9. "GSK Marketing" shall mean GSK personnel responsible for marketing GSK Products.
10. "GSK Medical Affairs" shall mean the organization within GSK consisting of highly trained experts with specialized scientific and medical knowledge, usually with an advanced scientific degree (e.g., an MD, PhD, or PharmD), whose role is limited to the provision of specialized, medical or scientific information, scientific analysis and/or scientific information to HCPs but excludes anyone performing sales, marketing, Promotional ride alongs, or other primarily commercial roles.
11. "GSK Product" or "GSK Products" shall mean: (1) Advair®; (2) Paxil®; (3)

Wellbutrin®; (4) any pharmaceutical or biological product approved by the Food and Drug Administration for the treatment of major depressive disorder; (5) any selective serotonin reuptake inhibitor (SSRI); and (6) any norepinephrine dopamine reuptake inhibitor (NDRI), that GSK Promotes or for which it directs Promotion.

12. “GSK Sales” shall mean the GSK sales force responsible for selling GSK Products.

13. “GSK Scientifically Trained Personnel” shall mean GSK personnel who are highly trained experts with specialized scientific and medical knowledge, usually with an advanced scientific degree (e.g., an MD, PhD, or PharmD), whose roles involve the provision of specialized, medical or scientific information, scientific analysis and/or scientific information to HCPs but excludes anyone performing sales, marketing, Promotional ride alongs, or other primarily commercial roles.

14. “Health Care Professional” or “HCP” shall mean any physician or other health care practitioner who is licensed to provide health care services or to prescribe pharmaceutical products.

15. “Meta-analyses” shall mean formal analyses combining evidence from independent studies using appropriate statistical methods, but shall not include any such analyses conducted in connection with the preparation or submission of an Investigational New Drug Application (IND), New Drug Application (NDA), Supplemental New Drug Application (sNDA), Abbreviated New Drug Application, (ANDA), nor shall it include any such analyses conducted in connection with any other regulatory report required under the Food, Drug and Cosmetic Act, 21 U.S.C. § 301 *et seq.* (FDCA), or by the U.S. Food and Drug Administration (FDA) or other regulatory body, to the extent the content or submission of which is treated as non-public or confidential by the relevant agency.

16. “Multistate Executive Committee” shall mean the Attorneys General and their staff representing Arizona, Florida, Illinois, Maryland, Oregon, Pennsylvania, Tennessee, and Texas.

17. “Multistate Working Group” shall mean the Attorneys General and their staff representing Alabama, Arizona, Arkansas, California, Colorado, Connecticut, Delaware, the District of Columbia, Florida, Georgia¹, Hawaii², Idaho, Illinois, Indiana, Iowa, Kansas, Kentucky, Maine, Maryland, Massachusetts, Michigan, Minnesota, Missouri, Montana, Nebraska, Nevada, New Jersey, New Mexico, New York, North Carolina, North Dakota, Ohio, Oklahoma, Oregon, Pennsylvania, Rhode Island, South Dakota, Tennessee, Texas, Utah³, Vermont, Virginia, Washington, Wisconsin, and Wyoming.

18. “Off-Label” shall mean a non-FDA approved use.

19. “Parties” shall mean the North Carolina Attorney General and GSK.

20. “Promotional,” “Promoting,” or “Promote” shall mean representations about a GSK Product intended to influence sales of that product, including attempts to influence prescribing practices and utilization of a GSK Product, that would be deemed Promotional labeling or advertising under the FDCA or any regulation promulgated thereunder, or by the FDA, under the most current draft or final standard promulgated by the FDA or the most current draft or final

¹ With regard to Georgia, the Administrator of the Fair Business Practices Act, appointed pursuant to O.C.G.A. § 10-1-395, is statutorily authorized to undertake consumer protection functions for the State of Georgia. References to the “States,” “Parties,” or “Attorneys General,” with respect to Georgia, include the Administrator of the Fair Business Practices Act.

² Hawaii is being represented on this matter by its Office of Consumer Protection, an agency which is not part of the state Attorney General’s Office, but which is statutorily authorized to undertake consumer protection functions, including legal representation of the State of Hawaii. For simplicity, the entire group will be referred to as the “Attorneys General,” and such designation, as it includes Hawaii, refers to the Executive Director of the State of Hawaii Office of Consumer Protection.

³ The Utah Attorney General’s Office represents the Utah Division of Consumer Protection (Division), the state agency charged with enforcement of the Consumer Sales Practices Act, in this action, but is not a party itself. As to Utah, the definition of “Attorneys General” means the Utah Attorney General as counsel to the Division.

FDA Guidance for Industry.

21. “Promotional Materials” shall mean any item used to Promote any GSK Product.
22. “Relevant State Consumer Protection Statutes” shall mean the consumer protection laws under which the Attorneys General have conducted the investigation.⁴
23. “Reprints Containing Off-Label Information” shall mean articles or reprints from a Scientific or Medical Journal, as defined in 21 C.F.R. 99.3(j), or Reference Publication, as defined in 21 C.F.R. 99.3(i), describing an Off-Label use of a GSK Product.
24. “Unsolicited Request” shall mean a request for information regarding a GSK Product communicated to an agent of GSK that has not been prompted by GSK.

III. COMPLIANCE PROVISIONS

Promotional Activities

- A. GSK shall not make, or cause to be made, any written or oral claim that is false, misleading, or deceptive about any GSK Product.
- B. GSK shall not represent that any GSK Product has any sponsorship, approval, characteristics, ingredients, uses, benefits, quantities, or qualities that it does not have.
- C. GSK’s policies and procedures shall address compensation (including through salaries, bonuses, or other means) for GSK Sales and GSK Marketing. These policies and procedures shall: (1) be designed to ensure that financial incentives do not inappropriately motivate GSK Sales or GSK Marketing to engage in improper sales Promotion, sales and marketing of GSK Products; and (2) include mechanisms, where appropriate, to exclude from incentive compensation sales that may indicate Off-Label Promotion of GSK Products. GSK shall make

⁴ In North Carolina, the relevant state consumer protection statute is N.C. Gen. Stat. § 75-1.1 *et seq.*

reasonable efforts in good faith to seek contractual language with any third-party contractor of prescriber-facing sales personnel requiring that any such personnel contracted to Promote GSK Products will not be compensated based on territory/individual level sales goals. GSK represents that, prior to the Effective Date, it implemented a program in the United States to eliminate incentive compensation based on territory/individual level sales goals for prescriber-facing sales personnel (e.g., sales representatives) and their direct managers (Patient First Program). The Patient First Program is described in more detail in Attachment A. GSK shall continue its Patient First Program or a substantially equivalent program through March 1, 2019.

The following paragraphs D through F shall be effective for a period of eight years from the Effective Date of this Judgment.

D. GSK shall not make in a Promotional context a representation or suggestion, not approved or permitted for use in the labeling or under the FDCA, that a GSK Product is better, more effective, useful in a broader range of conditions or patients, safer, has fewer, or less incidence of, or less serious side effects or contraindications than has been demonstrated by substantial evidence, or substantial clinical experience (as described in paragraphs (e)(4)(ii)(b) and (c) of 21 C.F.R. § 202.1), whether or not such representations are made by comparison with other drugs or treatments, and whether or not such a representation or suggestion is made directly or through use of published or unpublished literature, quotations, or other references.

E. GSK shall not Promote any GSK Product by use of Promotional Materials that:

1. contain a drug comparison that represents or suggests that a drug is safer or more effective than another drug in some particular when it has not been demonstrated to be safer or more effective in such particular by substantial evidence or substantial clinical experience;

2. contain a representation or suggestion that a drug is safer than it has been demonstrated to be by substantial evidence or substantial clinical experience, by selective presentation of information from published articles or other references that report no side effects or minimal side effects with the drug or otherwise selects information from any source in a way that makes a drug appear to be safer than has been demonstrated;
3. present information from a study in a way that implies that the study represents larger or more general experience with the drug than it actually does; or
4. use statistics on numbers of patients or counts of favorable results or side effects, derived from pooling data from various insignificant or dissimilar studies in a way that suggests either that such statistics are valid if they are not or that they are derived from large or significant studies supporting favorable conclusions when such is not the case.

F. When presenting information about a clinical study regarding GSK Products in any Promotional Materials, GSK shall not do any of the following for information that may be material to an HCP prescribing decision:

1. present favorable information or conclusions from a study that is inadequate in design, scope, or conduct to furnish significant support for such information or conclusions;
2. use the concept of statistical significance to support a claim that has not been demonstrated to have clinical significance or validity, or fails to reveal the range of variations around the quoted average results; or
3. use statistical analyses and techniques on a retrospective basis to discover and cite

findings not soundly supported by the study, or to suggest scientific validity and rigor for data from studies the design or protocol of which are not amenable to formal statistical evaluations.

Clinical Research

The following subsection shall be effective for eight years from the Effective Date of this Judgment.

G. GSK shall report research in an accurate, objective, and balanced manner as follows and as required by applicable law. To the extent permitted by the National Library of Medicine and as required by the FDA Amendments Act of 2007 (Public Law No. 110-85), GSK shall register GSK-sponsored Applicable Clinical Trials beginning after the Effective Date with the applicable registry and submit results of GSK-sponsored Applicable Clinical Trials completed after the Effective Date to the registry and results data bank as required by the FDA Amendments Act and any accompanying regulations that may be promulgated pursuant to that Act.

H. When submitting a manuscript on the results of a clinical study regarding any GSK Product for publication, GSK shall:

1. adhere to the ICMJE Uniform Requirements for Manuscripts Submitted to Biomedical Journals: Writing and Editing for Biomedical Publications, including authorship criteria, unless the applicable journal or congress to which the publication is submitted has more stringent requirements, in which case the journal or congress criteria for authorship will be followed;
2. acknowledge GSK's role as a funding source of the study which is the subject of the manuscript; and
3. disclose any change to the plan for the statistical analysis for that clinical study if

such change is inconsistent with GSK's standard operating procedure for Development, Review and Approval of Reporting and Analysis Plans. GSK's standard operating procedure for Development, Review and Approval of Reporting and Analysis Plans shall include requirements that such plans shall be consistent with the study protocol and shall be finalized before the date of final database release or interim database release (for an unblinded interim analysis).

- I. For any GSK Product, GSK shall also post on GSK's clinical study registry any observational studies or Meta-analyses conducted by GSK that are designed to inform the effective, safe, and/or appropriate use of any GSK Product.

Product Sampling

The following subsection shall be effective for five years from the Effective Date of this Judgment.

- J. GSK shall not provide samples of GSK Products to those HCPs who are not expected to prescribe the sampled GSK Products for an approved use, but who would be expected to prescribe the sampled product for an Off-Label use.
- K. If an HCP who would not be expected to prescribe the GSK Product for an approved use, but who would be expected to prescribe the product for an unapproved use, requests samples of that GSK Product, GSK personnel shall refer the HCP to GSK Medical Affairs where the practitioner can speak directly with a GSK Medical Affairs representative who will provide answers to the HCP's questions about the GSK Product and GSK may provide him/her with samples only if appropriate (i.e., if the HCP requests the samples for an FDA approved ("on-label") use).

Reprints

The following subsection shall be effective for five years from the Effective Date of this Judgment.

L. GSK shall not disseminate information describing any Off-Label use of a GSK Product, unless such information and materials are consistent with applicable FDA regulations and FDA Guidances for Industry.

M. Reprints Containing Off-Label Information regarding a GSK Product:

1. shall be accompanied by the FDA-approved labeling for the product, or a clearly and conspicuously described hyperlink that will provide the reader with such information;
2. shall contain a disclosure that is prominently displayed, which would include the first page or as a cover page where practicable, indicating that the article discusses Off-Label information; and
3. shall not be referred to or used in a Promotional manner.

N. GSK shall not disseminate any Reprint Containing Off-Label Information that relates to studies submitted to the FDA that were reviewed and specifically rejected by the FDA.

O. Nothing in this Judgment shall preclude GSK from revising its policies and practices regarding the dissemination of Reprints Containing Off-Label Information to be consistent with applicable FDA regulations and FDA Guidances for Industry that are revised or newly issued after the Effective Date of this Judgment.

Clinical Responses

The following subsection shall be effective for five years from the Effective Date of this Judgment.

P. GSK, through GSK Scientifically Trained Personnel, shall have ultimate responsibility for developing and approving all Clinical Responses regarding a GSK Product, including any that may describe Off-Label information. Additional approvals may be provided by the GSK Law Department. GSK shall not distribute any such materials unless:

1. Clinically Relevant Information is included in these materials to provide scientific balance;
2. data in these materials are presented in an unbiased, non-Promotional manner; and
3. these materials are clearly and conspicuously distinguishable from sales aids and other Promotional Materials.

Nothing in this subsection shall prohibit GSK Scientifically Trained Personnel from disseminating materials that are permitted to be distributed under Federal law.

Q. GSK Sales and GSK Marketing personnel shall not develop the medical content of Clinical Responses regarding a GSK Product.

R. Clinical Responses regarding a GSK Product may be disseminated only by GSK Scientifically Trained Personnel to HCPs, and GSK Sales and GSK Marketing personnel shall not disseminate these materials to HCPs except in circumstances implicating public health and safety issues. In such circumstances, GSK Sales and GSK Marketing personnel may disseminate a Clinical Response directly to HCPs when expressly authorized by the Health Care Compliance Officer, the Vice President of Medical/Scientific Affairs responsible for the GSK Product(s) included in the Clinical Response(s), and counsel from the GSK Law Department.

Responses to Unsolicited Requests for Off-Label Information

The following subsection shall be effective for five years from the Effective Date of this Judgment.

S. In responding to an Unsolicited Request for Off-Label information regarding a GSK Product, including any request for a specific article related to Off-Label uses, GSK shall:

1. advise the requestor that the request concerns an Off-Label use; and
2. inform the requestor of the drug's FDA-approved indication(s), provide labeling information and, where relevant to the Unsolicited Request, provide dosage information.

T. If GSK elects to respond to an Unsolicited Request for Off-Label information regarding a GSK Product, GSK Scientifically Trained Personnel shall provide specific, accurate, objective, and scientifically-balanced responses. Any such response shall not Promote a GSK Product for any Off-Label use(s).

U. Any written response to an Unsolicited Request for Off-Label information regarding a GSK Product shall include:

1. an existing Clinical Response prepared in accordance with Section III.P-R.
2. a Clinical Response prepared in response to the request in accordance with Section III.P-R; or
3. a report containing the results of a reasonable literature search using terms from the request.

V. Only GSK Scientifically Trained Personnel may respond in writing to an Unsolicited Request for Off-Label information regarding a GSK Product.

W. GSK Sales and GSK Marketing personnel may respond orally to an Unsolicited Request for Off-Label information regarding a GSK Product only by offering to request on behalf of the requester that a Clinical Response prepared in accordance with Section III.P-R or other information set forth in the current section above be sent in follow-up or by offering to put the

requester in touch with GSK Medical Affairs. GSK Non-Scientifically Trained Personnel shall not characterize, describe, identify, name, or offer any opinions about or summarize any such Off-Label information.

Grants

The following subsection shall be effective for five years from the Effective Date of this Judgment.

- X. GSK shall disclose information about medical education grants, including continuing medical education (“CME”) grants, regarding a GSK Product as required by applicable law.
- Y. GSK Medical Affairs shall manage all requests for funding related to medical education grants relating to a GSK Product. Approval decisions shall be made by GSK Medical Affairs, and shall be kept separate from the GSK Sales and GSK Marketing organizations.
- Z. GSK shall not use medical education grants or any other type of grant to Promote a GSK Product. This provision includes, but is not limited to, the following prohibitions:
 - 1. GSK Sales and GSK Marketing personnel shall not initiate, coordinate or implement grant applications on behalf of any customer or HCP;
 - 2. GSK Sales and GSK Marketing personnel shall not be involved in selecting grantees or medical education speakers; and
 - 3. GSK shall not measure or attempt to track in any way the impact of grants or speaking fees on participating HCPs’ subsequent prescribing habits, practices or patterns.
- AA. GSK shall not condition funding of a medical education program grant request relating to a GSK Product upon the requester’s selection or rejection of particular speakers.
- BB. GSK shall not suggest, control, or attempt to influence the specific topic, title, content,

speakers or audience for CMEs relating to a GSK Product, consistent with Accreditation Council for Continuing Medical Education (ACCME) guidelines.

CC. GSK Sales and GSK Marketing personnel shall not approve grant requests regarding a GSK Product, nor attempt to influence the awarding of grants to any customers or HCPs for their prescribing habits, practices, or patterns.

DD. GSK shall contractually require each medical education provider to clearly and conspicuously disclose to attendees of a medical education program regarding any GSK Product(s) GSK's financial support of the medical education program and any financial relationship with faculty and speakers at such medical education program.

EE. After initial delivery of a CME program regarding a GSK Product, GSK shall not knowingly fund the same program, nor shall it provide additional funding for re-distribution of the same program, if the program's speakers are Promoting a GSK Product for Off-Label use in that program.

IV. DISBURSEMENT OF PAYMENTS: PAYMENT TO THE STATES

Within 30 days of the Effective Date of this Consent Judgment, GSK shall pay \$105 million to be divided and paid by GSK directly to each Attorney General of the Multistate Working Group in an amount designated by and in the sole discretion of the Multistate Executive Committee.⁵ Said payment to North Carolina shall be used by the North Carolina Attorney General for attorneys' fees and other costs of investigation and litigation, or to be placed in, or applied to, the consumer protection enforcement fund, consumer education or litigation or local consumer aid or revolving fund, used to defray the costs of the inquiry leading hereto, or for

⁵ The State of North Carolina's share is two million five-hundred ninety-seven thousand eight-hundred thirty nine dollars and thirty three cents (\$2,597,839.33).

other uses permitted by state law, at the sole discretion of the North Carolina Attorney General. The Parties acknowledge that the payment described herein is not a fine or penalty, or payment in lieu thereof.

V. REPRESENTATIONS AND WARRANTIES

A. GlaxoSmithKline acknowledges that it is a proper party to this Consent Judgment. GlaxoSmithKline further warrants and represents that the individual signing this Consent Judgment on behalf of GlaxoSmithKline is doing so in his or her official capacity and is fully authorized by GlaxoSmithKline to enter into this Consent Judgment and to legally bind GlaxoSmithKline to all of the terms and conditions of the Consent Judgment.

B. The Attorney General warrants and represents that he is signing this Consent Judgment in his official capacity, and that he is fully authorized by his State to enter into this Judgment, including, but not limited to, the authority to grant the release contained in Section VI of this Consent Judgment, and to legally bind his State to all of the terms and conditions of this Consent Judgment.

VI. RELEASE

A. By execution of this Consent Judgment, the State of North Carolina releases and forever discharges GSK and all of its past and present, assigns, directors, divisions, employees, officers, parents, predecessors, shareholders, subsidiaries, successors, and transferees (collectively, the "Released Parties"), from the following: all civil claims, causes of action, parens patriae claims, damages, restitution, fines, costs, attorneys' fees, remedies and/or penalties that were or could have been asserted against the Released Parties by the Attorney General under N.C. Gen. Stat. § 75-1.1 *et seq.* or any amendments thereto, or by common law claims concerning unfair, deceptive, or fraudulent trade practices resulting from the Covered Conduct, up to and including

the Effective Date of this Consent Judgment (collectively, the “Released Claims”).

B. Notwithstanding any term of this Consent Judgment, specifically reserved and excluded from the Released Claims as to any entity or person, including Released Parties, are any and all of the following:

1. Any criminal liability that any person or entity, including Released Parties, has or may have to the State of North Carolina;
2. Any civil or administrative liability that any person or entity, including Released Parties, has or may have to the State of North Carolina, under any statute, regulation, or rule not expressly covered by the release in Section VI.A including, but not limited to, any and all of the following claims:
 - a. State or federal antitrust violations;
 - b. Medicaid violations, including, but not limited to, federal Medicaid drug rebate statute violations, Medicaid fraud or abuse, and/or kickback violations related to North Carolina’s Medicaid program;
 - c. Claims involving “best price,” “average wholesale price,” or “wholesale acquisition cost;”
 - d. State false claims violations; and
 - e. Claims to enforce the terms and conditions of this Consent Judgment.
3. Actions of state program payors of the State of North Carolina arising from the Covered Conduct, except for the release of civil penalties under the Relevant State Consumer Protection Laws.
4. Any claims individual consumers have or may have under the State of North Carolina’s consumer protection laws against any person or entity, including

Released Parties.

VII. CONFLICTS

If, subsequent to the Effective Date of this Consent Judgment, the federal government or any state, or any federal or state agency, enacts or promulgates legislation or regulations with respect to matters governed by this Consent Judgment that creates a conflict with any provision of the Consent Judgment and GSK intends to comply with the newly enacted legislation or regulation, GSK shall notify the Attorneys General (or the Attorney General of the affected State) of the same. If the Attorney General agrees, he/she shall consent to a modification of such provision of the Consent Judgment to the extent necessary to eliminate such conflict. If the Attorney General disagrees and the Parties are not able to resolve the disagreement, GSK shall seek a modification from an appropriate court of any provision of this Consent Judgment that presents a conflict with any such federal or state law or regulation. Changes in federal or state laws or regulations, with respect to the matters governed by this Consent Judgment, shall not be deemed to create a conflict with a provision of this Consent Judgment unless GSK cannot reasonably comply with both such law or regulation and the applicable provision of this Consent Judgment.

VIII. DISPUTE RESOLUTION

A. For the purposes of resolving disputes with respect to compliance with this Consent Judgment, should any of the signatory Attorneys General believe that GSK has violated a provision of this Consent Judgment subsequent to the Effective Date, then such Attorney General shall notify GSK in writing of the specific objection, identify with particularity the provisions of this Consent Judgment that the practice appears to violate, and give GSK 30 days to respond to the notification.

B. Upon receipt of written notice from any of the Attorneys General, GSK shall provide a good-faith written response to the Attorney General notification, containing either a statement explaining why GSK believes it is in compliance with the Consent Judgment or a detailed explanation of how the alleged violation occurred and statement explaining how and when GSK intends to remedy the alleged violation.

C. Except as set forth in Sections VIII.E and F below, the Attorney General may not take any action during the 30 day response period. Nothing shall prevent the Attorney General from agreeing in writing to provide GSK with additional time beyond the 30 days to respond to the notice.

D. The Attorney General may not take any action during which a modification request is pending before a court pursuant to Section VII, except as provided for in Sections VIII.E and F below.

E. Nothing in this Consent Judgment shall be interpreted to limit the State's Civil Investigative Demand (CID) or investigative subpoena authority.

F. The Attorney General may assert any claim that GSK has violated this Consent Judgment in a separate civil action to enforce compliance with this Consent Judgment, or may seek any other relief afforded by law, but only after providing GSK an opportunity to respond to the notification as described above and to remedy the alleged violation within the 30-day response period as described above, or within any other period as agreed to by GSK and the Attorney General; provided, however, that the Attorney General may take any action if the Attorney General believes that, because of the specific practice, a threat to the health or safety of the public requires immediate action.

IX. COMPLIANCE WITH ALL LAWS

Except as expressly provided in this Consent Judgment, nothing in this Consent Judgment shall be construed as:

1. relieving GSK of its obligation to comply with all applicable state laws, regulations, or rules, or granting permission to engage in any acts or practices prohibited by any law, regulation, or rule; or
2. limiting or expanding in any way any right any state represented by the Multistate Working Group may otherwise have to enforce applicable state law or obtain information, documents, or testimony from GSK pursuant to any applicable state law, regulation, or rule, or any right GSK may otherwise have to oppose any subpoena, civil investigative demand, motion, or other procedure issued, served, filed, or otherwise employed by the State pursuant to any such state law, regulation, or rule.

X. GENERAL PROVISIONS

A. Nothing in this Consent Judgment is intended to modify any prior settlement agreements between North Carolina and GlaxoSmithKline LLC formerly known as SmithKline Beecham Corporation, d/b/a GlaxoSmithKline, and SB Pharmco Puerto Rico, Inc.

B. Nothing will prevent the Attorney General from agreeing in writing to provide GSK with additional time to perform any act required by the Consent Judgment. The Attorney General shall not unreasonably withhold his/her consent to the request for additional time.

C. To the extent that any provision of this Consent Judgment obligates GSK to change any policy(ies) or procedure(s) and to the extent not already accomplished, GSK shall implement the policy(ies) or procedure(s) as soon as reasonably practicable, but no later than 120 days after the

Effective Date.

D. All notices under this Consent Judgment shall be sent by overnight United States mail.

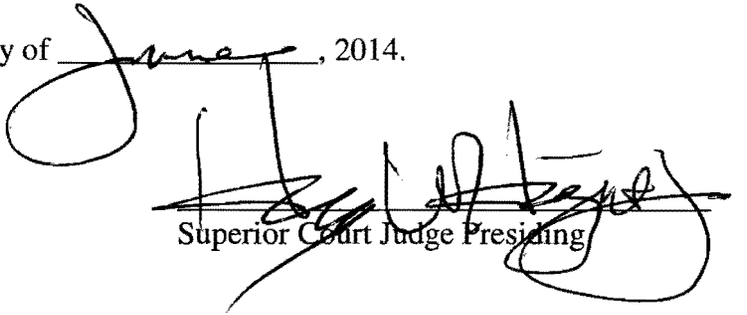
The documents shall be sent to the following addresses:

For GlaxoSmithKline LLC:

Matthew J. O'Connor
Covington & Burling LLP
1201 Pennsylvania Avenue, NW
Washington, DC 20004-2401

For the State of North Carolina:
Stuart M. Saunders
Assistant Attorney General
Consumer Protection Division
North Carolina Department of Justice
114 W. Edenton St.
Raleigh, NC 27603
Tel: 919-716-6031
Fax: 919-716-6050
ssaunders@ncdoj.gov

SO ORDERED, this the 4th day of January, 2014.


Superior Court Judge Presiding

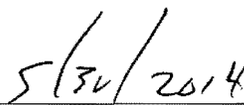
JOINTLY APPROVED AND
SUBMITTED FOR ENTRY:

FOR DEFENDANT GLAXOSMITHKLINE LLC

By:



Geoffrey E. Hobart
Matthew J. O'Connor
Covington & Burling LLP
1201 Pennsylvania Avenue, NW
Washington, DC 20004-2401



Date

FOR DEFENDANT GLAXOSMITHKLINE LLC

By: Frederick W. Rom

Frederick W. Rom

North Carolina Bar No. 26675

Womble Carlyle Sandridge & Rice, PLLC

150 Fayetteville Street, Suite 2100

Raleigh, NC 27601

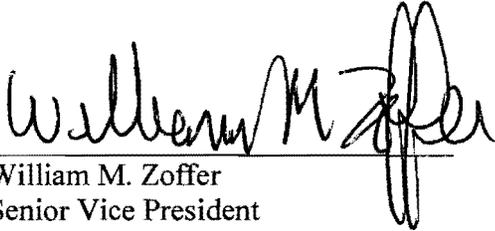
Local Counsel for GlaxoSmithKline LLC

5/23/14
Date

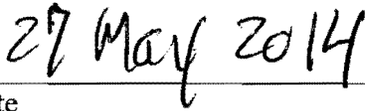
APPROVED:

FOR DEFENDANT GLAXOSMITHKLINE LLC

By:



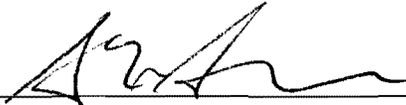
William M. Zoffer
Senior Vice President
GlaxoSmithKline LLC



Date

FOR PLAINTIFF, STATE OF NORTH CAROLINA

ROY COOPER
BY:



Stuart M. Saunders
Assistant Attorney General
Consumer Protection Division
N.C. Department of Justice
114 W. Edenton St.
Raleigh, N.C. 27603
Tel: (919) 716-6032
Fax: (919) 716-6050
ssaunders@ncdoj.gov

Date

6/2/14

ATTACHMENT A

Employee and Executive Incentive Compensation Policies and Practices

Pursuant to its existing Patient First Program, GSK agrees that it will not provide financial reward (through compensation, including incentive compensation or otherwise) or discipline (through tangible employment action) its prescribing-customer-facing field sales professionals (pharmaceutical sales representatives) or their direct managers based upon the volume of sales of GSK Products within a given employee's own territory or the manager's district. The Patient First Program includes evaluations for sales representatives based on business acumen, customer engagement, and scientific knowledge about GSK's Products. GSK shall continue its Patient First Program, or a substantially equivalent program through March 1, 2019. GSK commits to maintaining through at least March 1, 2019, absent agreement otherwise with the Multistate Executive Committee, the restrictions on such tangible employment decisions set forth in its Use of Territory/Individual Sales Data policy.