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1 Purpose

The Non-GMO Project (“the Project”) is a 501(c)(3) non-profit organization, incorporated in January 2007.

1.1 Mission

The Non-GMO Project is a non-profit organization offering rigorous product verification and trustworthy education that empowers people to care for themselves, the planet, and future generations.

1.2 Vision

A world in which humans respect and care for the community of life, making empowered choices that serve collective well-being and ecological harmony.

1.3 Guiding Beliefs

The work of the Project is guided by the following beliefs:

1. Everyone has a right to know what is in their food and deserves access to non-GMO choices.
2. By voting with our dollars every time we shop, collectively we have the power to change the way our food is grown and made.
3. Preserving and building the non-GMO supply chain is a critical step of transitioning toward a safe, healthy food supply for future generations.
4. The integrity of our diverse genetic inheritance is essential to human and environmental health and ecological harmony.
5. By encouraging a non-GMO seed supply, we are supporting the restoration of traditional seed breeding and the right of farmers to save and plant their own seeds and grow varieties of their choice.
6. A verified non-GMO system supports organic agriculture by reducing contamination pressure and protecting the supply of non-GMO seed.

1.4 Theories of Change

In fulfilling our mission, the Non-GMO Project operates according to the following theories of change:

1. Ensuring access to non-GMO choices necessitates the conversion of North American acreage to non-GMO agriculture at sufficient scale.
2. Efficient supply chain transformation requires consistent, uniform standards.
3. Standards must be both meaningful and achievable; meaningfulness includes valid testing and adherence to international protocols for credible standard setting.

4. Change of the magnitude proposed in the Non-GMO Project’s mission and vision depends on diplomatic, pragmatic and strategic cooperation with a diverse spectrum of stakeholders.

2 Stakeholder Engagement and Representation

2.1 Purpose

The Non-GMO Project strives to create consensus and action among a broad and diverse base of stakeholder interests. As part of its charitable 501(c)(3) mission, the Project also seeks to educate the public about the GMO issue and the value of non-GMO options.

2.2 Structure

The Project categorizes stakeholder interests into the following eight areas:

1. Retailers and Distributors
2. Input and Ingredient Processors
3. Consumer Packaged Goods Manufacturers
4. Livestock producers
5. Crop producers
6. Seed producers
7. Research and Policy
8. Consumer

2.3 Procedures

The Non-GMO Project engages stakeholders through a variety of activities and media, including but not limited to conferences, trade shows, educational and training events, newsletters, email, the Project’s own website, and social media applications such as Facebook, Twitter, and LinkedIn. Through these varied avenues, the Project educates its stakeholders about the GMO issue, informs them about its position and activities, and affords interested parties the opportunity to interact with the Project and provide feedback.

One key focus of stakeholder engagement is the solicitation of input during public comment periods. For details on this process, see section 7.1 of these terms of reference.

The Project is committed to growing its stakeholder base across all of the aforementioned stakeholder categories to assure broad and deep support for the Project’s programs. These
stakeholder categories are reflected throughout the governance and decision-making structures of the Project.

3  Board of Directors

3.1  Purpose
The Board of Directors (“Board”) exists to govern the Project, holding final decision-making authority and legal responsibility for the organization.

3.2  Structure
The Board shall consist of at least seven (7) but no more than thirteen (13) directors. The overall composition of the Board shall have a balanced representation from both large and small organizations.

3.3  Appointment Process
The directors shall be elected and/or reappointed at a regular annual meeting of the Board, typically in November of each year. The Board shall seek consensus on the election or reappointment of any candidate, per section 3.5, below. In cases where consensus cannot be achieved, election or reappointment shall be decided by majority vote.

3.4  Term
Directors are elected to a term of three (3) years. Each director, including a director elected to fill a vacancy, shall hold office for a maximum of three (3) terms, or a total of nine (9) years, whichever is longer, except as otherwise provided by law.

3.5  Decisions
The Board shall strive for consensus in its decisions. Consensus is defined as a lack of sustained opposition. Directors expressing a minority position shall be compelled to propose alternative solutions to resolve any given issue where consensus does not yet exist. In any given meeting of the Board if consensus cannot be reached the issue will be tabled until a subsequent meeting, during the interim of which the dissenting Director(s) shall be responsible for working with the project’s Executive Director and other members of the Board to achieve resolution.

If consensus is still not achievable at the subsequent meeting of the Board, the act of a majority of the directors present at a meeting duly held at which a quorum is present shall be the act of
the Board. Dissenting opinions shall be duly noted in the meeting minutes, if so desired by the dissenting party(ies).

### 3.6 Meetings & Quorum

The Board generally meets on approximately a bi-monthly basis. Annually, two of these meetings are in-person retreats (one day in length, preceded by a dinner meeting) and three of these meetings are held via conference (two hours in length).

A majority of directors shall constitute a quorum for the transaction of business at any meeting of the Board.

Proxies are not allowed, but guests may participate in meetings of the Board, by invitation of any Board member or by the Executive Director of the Project. All guests require the approval of the Board Chair.

In certain cases, voting may be conducted by written consent (i.e. email voting), unless any director expresses a need for limited additional discussion.

The Executive Director shall ensure that written minutes for all meetings are distributed to all directors via email following each meeting (typically within two weeks).

### 3.7 Committees

Each director shall serve on one of three standing committees, each of which generally meets on a monthly basis, or as outlined in the Annual Board Calendar. Each Committee shall be chaired by one director who is responsible for working with Project staff to set agendas, run meetings, and manage committee work.

#### 3.7.1 Executive Committee

The Executive Committee (“EC”) is chaired by the Board Chair and is comprised of at least three directors. The EC is responsible for compliance with certain fiduciary responsibilities, including:

1. Corporate filings (as detailed on the Board’s Google site)
2. Management of the Executive Director (including annual performance review and compensation)
3. Compliance with Terms of Reference and bylaws

#### 3.7.2 Governance Committee

The Governance Committee (“GC”) is comprised of at least three directors. The GC is responsible for Board effectiveness, including:

1. Board succession, including nominating and onboarding
2. Board policies
3. Board development (retreat agendas, trainings, annual Board evaluation etc.)

3.7.3 Standards Oversight Committee
The Standards Oversight Committee (“SOC”) is comprised of at least three directors. The SOC is responsible for ensuring the ongoing technical integrity of the organization’s Product Verification Program, including:

1. Governance of the Standards Committee (“SC”), including the development and implementation of strategies for: 1) Recruitment, 2) Orientation and Onboarding, 3) Management and Development, and 4) Succession
2. Adherence of the SC to the organization’s Guiding Statements
3. Adherence of SC members with the Code of Conduct
4. Facilitation of conflict resolution within the SC, as appropriate
5. Approval of and adherence with the Standards Revision Process
6. Ratification of the Standard in consultation with the Board or on behalf of the Board, as appropriate

3.7.4 Finance Committee
The Finance (“FC”) is chaired by the Treasurer and is comprised of at least three directors. The FC is responsible for the financial health of the organization, including:

1. Annual Budget
2. Quarterly financial statements
3. Cash flow
4. Accurate and timely tax filing
5. Investments, short term and long term for organizational health

3.7.5 Audit Committee
The Audit Committee (“AC”) is comprised of at least two directors. The AC is responsible for the overseeing the annual financial audit, internal controls and risk management of the organization, including oversight of:

1. Accounting practices
2. Annual audits and reviews
3. External auditor selection
4. Accurate and timely tax filing
5. Risk management practices
3.8 Code of Conduct
All directors must agree to abide by the Code of Conduct set forth in Appendix A of these terms of reference.

3.9 Attendance
Two consecutive absences from biannual retreats, or three consecutive absences from any scheduled meetings shall be considered resignation from the Board unless the Executive Committee makes an exception for extenuating circumstances.

3.10 Officer Roles and Responsibilities
Board officers oversee and direct major aspects of organization governance. Officers are elected annually and include a Board Chair, Vice Chair, Secretary and Treasurer. Responsibilities of each role include:

3.10.1 Board Chair
1. Provide leadership for the Board in performing its duties
2. Provide oversight and guidance to the Executive Director
3. Plan meetings of the Board; prepare agendas with Executive Director
4. Chair Board meetings and Executive Committee meetings
5. Coordinate Board level annual planning process with Executive Director
6. Ensure an established process for Executive Director monitoring, evaluation and compensation reviews; including clearly articulated goals and expectations from the Board to the Executive Director
7. Ensure new Board member orientation process is clear and consistently executed

3.10.2 Vice Chair
1. Understand the basic functions associated with Board Chair responsibilities
2. Step into the role of the Board Chair when needed
3. Coordinate annual Board succession process with Board Chair
4. Serve on the Executive Committee

3.10.3 Secretary
1. Ensure timely notice of meetings, and of agendas of the Board meetings
2. Ensure legal files are maintained for minutes of the Board and Committees
3. Represent the organization on legal documents when an officer signature is required
4. Serve on the Executive Committee
3.10.4 Treasurer

1. Chair Finance & Audit Committee meetings
2. Provide quarterly fiscal oversight to the financial health of the Project
3. Ensure annual budget process is in place, executed and prepared for Board action
4. Oversee the annual audit process from auditor selection through Board reporting
5. Monitor and oversee bank accounts and financial investments
6. Ensure that all required tax filings and payments meet IRS and State regulation

4 The Standard

4.1 Purpose

The Non-GMO Project’s Standard aims to verify that systems are in place for:

1. **Testing**: Meaningful, ongoing testing of high GMO risk inputs.
2. **Traceability**: Supply chain traceability, especially following input testing.
3. **Segregation**: Protecting compliant inputs from contamination by non-compliant inputs.
4. **Formulation**: Obtaining inputs in accordance with uniform and meaningful specifications.
5. **Labeling**: Accurate and clear product labeling.
6. **Quality assurance**: Maintaining operational consistency and addressing non-conformities promptly.

4.2 Requirements

The full Non-GMO Project Standard is available on the Project’s website:

[www.nongmoproject.org](http://www.nongmoproject.org)

4.3 Governance

The Standard is maintained by a Standards Committee, described in section 6 of these terms of reference, along with oversight by the Board Standards Oversight Committee, described in section 3.7 of these terms of reference.

At the discretion of the Board, the Project may form special Technical Advisory Committees to inform development on particular aspects of the Standard. Such committees shall operate according to terms outlined by the Board and/or staff, and shall be publicly listed on the Project’s website.
4.4 Interpretation
Interpretation of the Standard is issued as needed in the form of TA Guidance, described in section 5.4, point 2, of these terms of reference.

4.5 Revisions
The process for updates to the Standard is described in section 7 of these terms of reference.

5 Product Verification Program

5.1 Purpose
The Project oversees a Product Verification Program (“PVP” or “Program”), the purpose of which is to assess products’ compliance with the Non-GMO Project Standard.

5.2 Methodology & Approach
1. The PVP is based on a practice/process-oriented Standard that uses testing as a key strategic tool to confirm that practices/processes are meeting expectations.
2. A core goal of the Project is to identify, create, and/or maintain sources and practices that effectively minimize GMO risk to the supply chain.
3. Release of products to the marketplace shall be contingent on products meeting requirements regarding Non-GMO Project Standard compliance, including traceability, segregation and testing.
4. Continuous improvement on the part of PVP Participants is required with the common goal of completely eliminating any GMO risk ingredients from the production chain.

5.3 Definitions
2. License Agreement (“LA”). Contract between the Project and a Participant outlining terms with regard to use of the Project’s trademarks.
3. Participant. A company that is seeking verification within the PVP and signs a License Agreement with the Project.
5. Product. A unique branded formula and process, where process could be either the manufacturing or facility process.
6. **Prospect**: A company that is a prospective Participant.

7. **Technical Administrator** ("TA"). An entity contracted to oversee evaluation of products for purposes of determining compliance with the Standard.

8. **Technical Administrator Portal** ("TAP"). The online platform used for coordinating TA communications, including TA Guidance (see 5.4, point 2, below).

9. **Trademarks**. Any of the Project’s registered trademarks; for details see section 8 of these terms of reference.

### 5.4 Technical Administrator Oversight

Ensuring quality and consistency in TA performance is a key focus of the Non-GMO Project.

1. **Appointment and scope.** The Project selects and approves TAs according to a rigorous process duly established by the Board. This process shall include an initial assessment of the TA’s demonstrated proficiency as a certifier for other programs as well as thorough technical training by Project staff. TA contracts are offered at the Board’s discretion and contain provisions including, but not limited to, administration of the PVP, data access, surveillance testing, collection of fees from Participants, reporting, quality of service, confidentiality and quality control.

2. **TA Guidance.** A central process in support of TA consistency is the issuance of TA Guidance related to interpretation of the Standard. TA Guidance is issued at the discretion of Non-GMO Project staff through a process that may include input from the Board and/or Standards Committee. Formal TA Guidance is issued to all TAs simultaneously, and is transparently maintained on the online Technical Administrator Portal.

### 5.5 Quality Assurance

The Project operates a variety of Quality Assurance ("QA") programs designed to ensure the technical integrity of the PVP. These programs are housed within the Project’s Standards & Verification Department.

1. **Surveillance Testing.** Designed to confirm the compliance of verified products with the relevant Action Threshold, as set forth in the Standard. Scope includes quarterly spot testing of randomly selected products with an emphasis on High-Risk and Monitored-Risk products and inputs.

2. **Product Auditing.** Designed to confirm the accuracy of TA decisions with regard to compliance with the Standard. Scope includes randomly selected products as well as products for which information presented to Project staff indicates potential cause for concern.
3. **Laboratory Oversight.** Designed to confirm satisfactory and equivalent competency across all NGP-approved laboratories. Scope includes an initial evaluation and annual renewal process based on compliance with requirements including, but not limited to, ISO 17025 accreditation and satisfactory proficiency testing (ring trial) results. A list of approved labs is maintained on the Project’s website: **www.nongmoproject.org**

5.6 **Database Management**

As part of its oversight of the PVP, the Project maintains a database of verified products and Participants. This database is used to support a number of critical Program operations, including populating public listings on the Project’s website and mobile phone apps.

5.7 **Customer Service**

The Project strives to provide excellent customer service to all Participants and Prospects. Accordingly, the Project’s Client Services Department supports brand owners throughout the verification process with educational information and assistance with complaint resolution.

6. **Standards Committee**

6.1 **Purpose**

The Standards Committee (“SC”) is a standing committee that exists to oversee the development of the Non-GMO Project Standard. Revisions and interpretations proposed by the SC are reviewed and decided upon by the SOC on behalf of the Board, which holds final decision-making authority. The SC is responsible for addressing all comments received from stakeholders with respect to the content and proposed revisions to the Standard.

The SC shall abide by the schedule and terms set forth by the Non-GMO Project.

6.2 **Structure & Qualifications**

SC members are independent of the SOC and Board and cannot serve on either while serving on the SC. Membership is open to all interested parties with approval of the SOC.

The SC shall consist of voting members and non-voting members.

Voting members shall comprise the voting body of the SC and shall consist of at least five (5) but no more than seven (7) members. No more than one (1) representative from the same company or organization may serve as a voting SC member at any given time.
Non-voting members shall comprise the non-voting body of the SC and may consist of one (1) representative from each TA at the discretion of the SOC, and Non-GMO Project staff as appropriate. TA representatives participate in a limited advisory capacity, providing information as requested by the voting SC members or the SOC.

Overall composition of the SC shall have a balanced representation of interests, including those outlined in section 2.2 of this document.

6.3 Appointment Process

The Non-GMO Project solicits interest through its website and other channels from persons wishing to serve on the SC. The SOC shall strive for consensus when appointing candidates to the SC. In cases where consensus cannot be achieved, appointment shall be decided by majority vote of the SOC.

6.4 Term

Voting SC members are elected to a seat and a term of three (3) years. Each voting SC member, including a SC member elected to fill a vacancy, shall hold office for a maximum of two (2) consecutive terms, or a total of six (6) years, whichever is shorter. After serving two (2) consecutive terms, former voting SC members may serve additional terms subject to these terms of reference after a gap of one (1) term.

The voting members of the SC shall be represented by no more than seven (7) seats. Seats are assigned term start and end dates. Terms are staggered such that no more than two (2) seats are assigned terms with the same end date.

Non-voting members shall serve 3-year terms and may be reappointed as appropriate at the discretion of the SOC.

6.5 Decisions

The SC strives for consensus when reaching conclusions to its deliberations. Consensus is defined as a lack of sustained opposition. Members expressing a minority position shall be compelled to propose alternative solutions to resolve any given issue where consensus does not yet exist.

If consensus is not achievable after reasonable effort, the act of a majority of the voting members present at a meeting duly held at which a quorum is present shall be the conclusion of the SC. In the event of a tie, the SOC Chair shall cast the deciding vote. Dissenting opinions shall be duly noted in the meeting minutes and brought forward to subsequent Board meetings by the SOC, if so desired by the dissenting party(ies).
6.6 Meetings & Quorum
The SC meets approximately quarterly, with more frequent meetings during Standards ratification cycles.

A majority of voting members shall constitute a quorum.

Proxies are not allowed, but guests may participate by approval of the SOC Chair.

In certain cases, voting may be conducted by written consent (i.e. email voting), unless any voting SC member expresses a need for limited additional discussion.

All SC meetings shall be open to Board members.

Written minutes shall be taken and shall at least detail any proposed changes to the Standard and any dissenting opinions in cases where consensus is not reached.

6.7 Code of Conduct
All SC members must agree to abide by the Code of Conduct set forth in Appendix A of these terms of reference.

6.8 Attendance
Three (3) consecutive absences from any scheduled meetings shall be considered resignation from the SC unless the SOC makes an exception for extenuating circumstances.

7 Revisions to The Non-GMO Project Standard
The Non-GMO Project Standard (“the Standard”) is intended to be a “living” document, capable of flexibly incorporating feedback from all stakeholders in such a way that it always maintains a viable balance of meaningfulness and achievability.

7.1 General Revisions and Public Comment Periods
The Project accepts comments from stakeholders at any time, through its website, and these are reviewed in an ongoing manner. Comments not addressed immediately shall be rolled into the next regularly scheduled process for revision of the Standard.

The Non-GMO Project Standard in its entirety is subject to a regular revision process. This revision process generally consists of two successive open public comment periods. The Project shall notify stakeholders of the start of the consultation periods via announcement on its website and by direct email to stakeholders for whom the Project has such contact data. The Project may, at its discretion, also make such announcements through other media.
In revision years, the first public comment period commences no later than April and lasts for 60 days, during which stakeholders may submit comments on the Standard through the Project’s website. Upon opening the public comment period, the Project may also specifically solicit stakeholders for feedback on particular questions about the Standard. During this comment period, any recommendations from special advisory committees shall also be considered.

At the close of the first round of public comment, the SC will review all comments received and propose corresponding changes to the Standard. These proposed changes, along with the SC’s rationale for the changes, will then be published for a second round of public comment, which shall be conducted in the same fashion as the first round. This second round will normally last for 60 days, but may be reduced to as little as 30 days if the proposed changes are not deemed by the SC to be contentious. In cases where there appears to be no new decisions that must be made following the initial 60-day consultation period (no areas of disagreement among stakeholders), the second comment period may be waived altogether. In any cases where the second round is shortened or cancelled, the rationale for this shall be explained on the Project’s website.

Following the second round, the SC will again review all comments received, and create a new final version of the Standard. Comments received in the second round will be addressed along with the SC’s response and rationale.

If after the second round there are still contentious issues, the Project may convene additional comment periods, each of which shall last no less than 30 days. The same procedures as for the earlier rounds shall be followed.

Once the SC has made a decision on a new final version of the Standard, the Standard shall be brought to the Board for final ratification. Although the Board is legally responsible for the Non-GMO Project, and therefore holds final decision-making authority, alignment between the SC and the Board is essential and shall be prioritized in the decision-making process. Full consensus of all SC and Board members shall be diligently sought prior to ratification of any new Standard.

In general, Participants must come into compliance with the revised Standard by time of their next renewal, but no sooner than 180 calendar days from the publication date. In certain cases, the Board may determine that additional time is needed for Participants to come into compliance with changes to the Standard. Any such extended timelines shall be incorporated in Appendix E of the published Standard as soon as practically possible. Along with the new version, the Project will also publish a summary of the changes made since the last version and a corresponding summary of the rationale for those changes. The Project shall also directly email the new Standard and the summary of changes to all stakeholders on its emailing list(s).
7.2 Special Revisions to the Non-GMO Project Standard

7.2.1 Revisions to Appendix B (List of High-Risk Inputs) and Appendix C (List of Monitored-Risk Inputs)

As a mechanism for maintaining accuracy in the Standard with regard to which crops and inputs are at risk for GMO contamination, the SC does have the ability to recommend changes to Appendices B and C (risk lists) between regular revision processes, and the Board has the authority to approve such changes between comment periods. Such revisions must adhere to the following guidelines:

a. A GM crop, input, or other organism shall be added to one of the risk lists if the GM event is:
   i. commercialized.
   OR
   ii. an unapproved event that is detected in the environment, food, or feed supply.

b. Similarly, a GM crop, input, or other organism may be removed from one of the GM risk lists if:
   i. Sufficient testing of relevant global risk areas demonstrates that the overall level of contamination with the GMO under consideration is consistently well below the relevant Non-GMO Project Standard Action Threshold.

7.2.2 Urgent Standard Revisions

Because the issue of GMOs is highly complex and often changes suddenly, the Project allows for urgent revisions to the Standard. These changes must be triggered by a majority vote of the Board, who may have received relevant input from the SC, a TA, the Executive Director, and/or other sources.

If the Board decides an urgent revision is warranted, the nature of the revision and any specific proposals related to the change will be made known to the SC, who shall have adequate opportunity to provide feedback. The Board and SC shall strive to reach consensus on any such proposal(s), and shall solicit additional input from stakeholders as needed to support an informed decision. The Board and SC shall abide by the decision-making rules established in this document.

Any decided changes to the Standard under this provision shall go into force upon their publication on the Project’s website and simultaneous notification of stakeholders by Project staff via direct email (plus any other media at the discretion of Project staff). As part of this notification the Project shall explicitly solicit stakeholder comment on the changes.
The changes made using this provision shall remain in force at least until the subsequent regular revision process.

8 Trademarks

A core asset of the Non-GMO Project is its trademarks, which include:

- Non-GMO Project Verified mark (English and Bilingual/French)
- Non-GMO Project logo
- Non-GMO Project name
- “Look for the Butterfly”

These trademarks are governed through Licensing Agreements with Participants, retailers, and other stakeholders at the Project’s sole discretion. One primary purpose of the trademarks is to make readily apparent to shoppers which products have successfully completed the Product Verification Program; see section 5 of these terms of reference.

The Project maintains a Trademark Use Guide to assist stakeholders in appropriate and legal use of the trademarks. The guide is available from the Project’s Marketing & Communications Team.
Appendix A: Code of Conduct

The Non-GMO Project has adopted the following Code of Conduct that all Board members and Standards Committee members must agree to.

A.1 Prohibition Against Private Inurement and Procedures for Managing Conflicts of Interest

No member of the Board of Directors (Board) or Standards Committee (SC) shall derive any personal profit or gain, directly or indirectly, by reason of his or her service as a Board or Standards Committee member with the Non-GMO Project. Members of the Board and SC shall conduct their personal affairs in such a manner as to avoid any possible conflict of interest with their duties and responsibilities as members of the Board or SC. Nevertheless, conflicts may arise from time to time.

1. When there is a decision to be made or an action to be approved that will result in a conflict between the best interests of the Non-GMO Project and the Board or SC member’s personal interests, the Board or SC member has a duty to immediately disclose the conflict of interest so that the rest of the Board’s or SC’s decision making will be informed about the conflict.

2. It is every Board and SC member’s obligation, in accordance with this policy, to ensure that decisions made by the Board and SC reflect independent thinking. Consequently, in the event that any Board or SC member receives compensation from the Non-GMO Project such compensation will be determined by and approved by the full Board in advance.

3. Any conflicts of interest, including, but not limited to financial interests, on the part of any Board or SC member, shall be disclosed to the Board when the matter that reflects a conflict of interest becomes a matter of Board or SC action, and through an annual procedure for all Board and SC members to disclose conflicts of interest.

4. Any Board or SC member having a conflict of interest shall not vote or use his or her personal influence to address the matter, and he or she shall not be counted in determining the quorum for the meeting.

5. All conflicts disclosed to the Board or SC will be made a matter of record in the minutes of the meeting in which the disclosure was made, which shall also note that the Board or SC member with a conflict abstained from the vote (and was not present for any discussion, as applicable) and was not included in the count for the quorum for that meeting.

6. Any new Board or SC member will be advised of this policy during board orientation and all Board and SC members will be reminded of the Code of Conduct and of the procedures for disclosure of conflicts and for managing conflicts on a regular basis, at least once a year.
7. This policy shall also apply to any Board and SC member’s immediate family or any person acting on his or her behalf.

A.2 Mission Alignment

Board and SC members must demonstrate strong, dedicated support for the Project’s mission of preserving and building sources of non-GMO products, educating consumers, and providing verified non-GMO choices. Accordingly, no Board or SC member may be an employee of nor knowingly maintain any direct interest, financial or otherwise, in organizations that explicitly develop or explicitly promote the use of GMOs.

A.3 Prohibition Against Sexual Harassment

The Non-GMO Project strives to maintain a workplace that is free from illegal discrimination and harassment. While all forms of harassment are prohibited, it is the organization’s policy to emphasize that sexual harassment is specifically prohibited. Any Board or SC member who engages in discriminatory or harassing conduct towards any individual is subject to removal from the Board or SC. Complaints alleging misconduct on the part of Board or SC members will be investigated promptly and as confidentially as possible by a task force of the Board appointed by the Executive Committee.

A.4 Confidentiality

Board and SC members are reminded that confidential financial, strategic, personnel and other matters concerning the organization, donors, staff or clients/consumers may be included in Board or SC materials or discussed from time to time. Board and SC members should not disclose such confidential information to anyone.

A.5 Participation

Board and SC members are expected to exercise the duties and responsibilities of their positions with integrity, collegiality, and care. This includes:

1. Making attendance at all meetings a high priority.
2. Being prepared to discuss the issues and business on the agenda, and having read all background material relevant to the topics at hand.
3. Actively engaging in discussions.
4. Cooperating with and respecting the opinions of fellow Board or SC members, and leaving personal prejudices out of all Board or SC discussions, as well as supporting actions of the Board or SC even when the Board or SC member personally did not support the action taken.
5. Putting the interests of the organization above personal interests.
6. Representing the organization in a positive and supportive manner at all times and in all places.
7. Refraining from publicly representing the Project, deferring instead to the Non-GMO Project’s appointed spokesperson(s).
8. Refraining from activities that pose a reputational risk to the Non-GMO Project (fraudulent claims, activities that could be construed as misleading or in contravention to the Non-GMO Project mission, etc.)
9. Showing respect and courteous conduct in all meetings.
10. Refraining from intruding on administrative issues that are the responsibility of management, except to monitor the results and ensure that procedures are consistent with policy.
Appendix B: NGP Org Chart