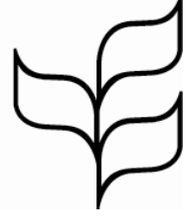


Distr.
GENERAL

UNEP/CBD/BS/COP-MOP/5/12
30 July 2010

ARABIC
ORIGINAL: ENGLISH

الاتفاقية المتعلقة بالتنوع البيولوجي



مؤتمر الأطراف في اتفاقية التنوع البيولوجي
العامل كاجتماع للأطراف في بروتوكول
قرطاجنة للسلامة البيولوجية
الاجتماع الخامس

ناغويا، اليابان، 11-15 أكتوبر/تشرين الأول 2010
البند 13 من جدول الأعمال المؤقت*

تقييم المخاطر وإدارة المخاطر (المادتان 15 و16)

مذكرة من الأمين التنفيذي

أولا - مقدمة

1. يحدد بروتوكول قرطاجنة للسلامة الأحيائية أحكاما تتعلق بتقييم المخاطر (المادة 15 والمرفق الثالث) لتحديد وتقييم التأثيرات المعاكسة المحتملة للكائنات المحورة الحية على صون التنوع البيولوجي واستخدامه. المستدام مع مراعاة المخاطر على صحة البشر، وإدارة المخاطر (المادة 16) لتمكين الأطراف من وضع ومراعاة الآليات والتدابير والاستراتيجيات الملائمة لتنظيم وإدارة ومكافحة المخاطر التي يتم التعرف عليها في عملية تقييم المخاطر وفقا لأحكام البروتوكول

2. وكان مؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول قد قرر، خلال اجتماعه الأول، أن ينظر في اجتماعه الخامس الطرق التي قد يمكن من خلالها تحديد هوية الكائنات المحورة الحية التي قد لا تنطوي على الأرجح على تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر وذلك بغرض التوصل إلى مقرر وفقا للفقرة 4 من المادة 7.¹

UNEP/CBD/BS/COP-MOP/5/1*.

¹ الفقرة 7 (أ) من المرفق بالمقرر BS-I/12.

3. وأتسأ مؤتمراً الأطراف، خلال اجتماعه الرابع، لدى النظر في مدى الحاجة إلى إرشاد آخر بشأن الجوانب النوعية لتقييم المخاطر وإدارة المخاطر، منتدى الكتروني مفتوح العضوية بشأن الجوانب النوعية لتقييم المخاطر من خلال غرفة تبادل معلومات السلامة الأحيائية وفريق الخبراء التقنيين المخصص المعني بتقييم المخاطر وإدارة المخاطر بالاختصاصات الواردة في مرفق المقرر. وعلاوة على ذلك، طلبت الأطراف في البروتوكول من الأمين التنفيذي أن يعقد (1) مجموعات مخصصة ومؤتمر الكتروني واحد على الأقل في الوقت الحقيقي لكل إقليم قبيل كل اجتماع لفريق الخبراء التقنيين المخصص المعني بتقييم المخاطر وإدارة المخاطر بغرض تحديد القضايا الرئيسية ذات الصلة بالجوانب النوعية لتقييم المخاطر وإدارة المخاطر على النحو المبين في المرفق بالمقرر، و(2) اجتماعين لفريق الخبراء التقنيين المخصص قبل الاجتماع الخامس لمؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول.²

4. وطلبت الأطراف أيضاً أثناء نظرها لبناء القدرات في مجال تقييم المخاطر، خلال اجتماعها الرابع من الأمين التنفيذي (أ) أن ينسق ويبصر، جنباً إلى جنب مع هيئات الأمم المتحدة الأخرى ذات الصلة وغيرها من المنظمات الدولية، إعداد وتدريب عن تقييم المخاطر وإدارة المخاطر فيما يتعلق بالكائنات المحورة الحية (2) عقد دورات تدريبية إقليمية أو دون إقليمية، قبل الاجتماع الخامس للأطراف، لتمكين البلدان من اكتساب خبرات مباشرة في إعداد وتقييم تقارير عمليات تقييم المخاطر وفقاً للبروتوكول و(3) عقد حلقة عمل بشأن بناء القدرات وتبادل الخبراء فيما يتعلق بتقييم المخاطر وإدارة المخاطر بخصوص الكائنات المحورة الحية في الإقليم الفرعي للمحيط الهادي.³

5. وعلاوة على معالجة مدى الحاجة إلى إرشاد آخر بشأن الجوانب النوعية لتقييم المخاطر على النحو المشار إليه في الفقرة 3، طلب من فريق الخبراء التقنيين المخصص أيضاً النظر في الطرائق المحتملة للتعاون في تحديد الكائنات المحورة الحية أو السلالات المحورة التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر التي تتعرض لها صحة البشر. وبغية مساعدة فريق الخبراء التقنيين المخصص في مداولاته، طلب مؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول من الأطراف ودعا الحكومات الأخرى والمنظمات ذات الصلة إلى تقديم المعلومات السليمة من الناحية العلمية المتوفرة بشأن تحديد هوية الكائنات المحورة الحية أو السلالات المحددة التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر. كما طلبت الأطراف من الأمين التنفيذي تجميع المعلومات المتلقاة وإعداد تقرير تجميعي للنظر من جانب فريق الخبراء التقنيين المخصص والأطراف.⁴

6. وعلى ذلك فإن هذه المذكرة أعدها الأمين التنفيذي لمساعدة الأطراف في البروتوكول في نظرها للبدء من جدول الأعمال المتعلق بتقييم المخاطر وإدارة المخاطر. ويتضمن القسم الثاني تحليلاً للنتائج الرئيسية لعملية وضع إرشاد آخر بشأن الجوانب النوعية لتقييم المخاطر. ويحتوي القسم الثالث على استعراض لأنشطة بناء القدرات التي اضطلع بها استجابة لاجتماع الأطراف. ويضم القسم الرابع عرضاً عاماً للتقديرات والتوصيات المتعلقة بالتعاون في تحديد هوية الكائنات المحورة الحية التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر⁵ ويقدم القسم الخامس بعض العناصر التي قد تساعد الأطراف في نظر طرائق تحديد الكائنات المحورة الحية التي قد لا يكون لها على الأرجح تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر⁶. ويستخلص القسم السادس بعض الاستنتاجات ويقترح بعض العناصر لمشروع مقرر للنظر من جانب الأطراف

ثانياً - الإرشاد الآخر بشأن الجوانب النوعية لتقييم المخاطر

7. سعياً إلى تنفيذ مختلف عناصر المقرر BS-IV/11 فيما يتعلق بوضع إرشاد آخر بشأن تقييم المخاطر، وضعت الأمانة، بالتشاور مع مكتب مؤتمر الأطراف العاملة كاجتماع للأطراف في البروتوكول عملية مستمرة تتألف من ثلاثة أنواع من الأنشطة (1) مجموعات نقاش إلكترونية مخصصة (2) مؤتمرات إقليمية إلكترونية في الوقت الحقيقي و(3) اجتماعات مباشرة لفريق الخبراء التقنيين المخصص.

8. وقد بدأت العملية بافتتاح منتدى الخبراء الإلكتروني مفتوح العضوية بشأن تقييم المخاطر وإدارة المخاطر (المنتدى الإلكتروني) من خلال غرفة تبادل معلومات السلامة الأحيائية.⁷

² الفقرات 3 و4 و6 من المقرر BS-IV/11.

³ الفقرتان 12 و13 من المقرر BS-IV/11

⁴ الفقرات 3 و4 و6 من المقرر BS-IV/11.

⁵ على النحو الوارد في الفقرة 4(ب) (3) من مرفق المقرر BS-I/12.

⁶ على النحو الوارد في الفقرة 7(أ) (1) من مرفق المقرر BS-I/12.

⁷ يتوافر على http://bch.cbd.int/onlineconferences/forum_RA.shtml.

9. ودعا الأمين التنفيذي، في إخطار، الأطراف والحكومات الأخرى والمنظمات ذات الصلة إلى ترشيح خبراء في تقييم المخاطر في المنتدى الإلكتروني باستخدام استمارة التعيين الموحدة لخبراء السلامة الأحيائية. واستعرضت الأمانة الترشيحات للتأكد من استكمالها وفقا للمعايير والمتطلبات الدنيا لخبراء السلامة الأحيائية على النحو الوارد في المقرر BS-IV/4.

10. وجرى تسجيل ما مجموعه 229 خبيراً في المنتدى الإلكتروني المفتوح العضوية. وكان من بينهم 53 ترشيحاً من عدد يبلغ 48 طرفاً و11 خبيراً من خمسة بلدان غير أطراف و65 خبيراً تم تسجيلهم كمراقبين.⁸

11. وكجزء من التحضير لعمل فريق الخبراء التقنيين المخصص، عقدت ثماني مجموعات نقاش إلكترونية مخصصة وأربعة مؤتمرات إلكترونية إقليمية في الوقت الحقيقي (أوروبا وأمريكا اللاتينية وأفريقيا وآسيا) في إطار المنتدى الإلكتروني فيما بين نوفمبر/ تشرين الثاني 2008 وفبراير/ شباط 2009.⁹

12. وقد جرى اختيار المشاركين في فريق الخبراء التقنيين المخصص على أساس مشاركتهم الفعالة في الأحداث الجارية في المنتدى الإلكتروني وفقاً لظروف العمل الموحدة للجهاز الفرعي المعنى بالمشورة العلمية والتقنية والتكنولوجية التابع لاتفاقية التنوع البيولوجي¹⁰ حسب الطلب الوارد في المقرر BS-IV/11 وبالتشاور مع مكتب مؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول. وترد قائمة المشاركين في فريق الخبراء التقنيين المخصص في المرفق الأول بهذه المذكرة.

13. وعقد الاجتماع الأول لفريق الخبراء التقنيين المخصص المعنى بتقييم المخاطر وإدارة المخاطر في مونتريال من 20 إلى 24 أبريل/ نيسان 2009. وحضر الاجتماع كأعضاء في الفريق ثماني عشر مشاركاً من سبع عشر طرفاً فضلاً عن ثمانية مراقبين من ثلاث بلدان غير أطراف وخمس منظمات.

14. ونفذ، خلال الاجتماعين اللذين عقدهما الفريق، عدد من الأنشطة بغرض تحقيق تقدم في إعداد مشروع إرشاد عن كل مسألة من المسائل النوعية التي حددت في الاجتماع الأول لفريق الخبراء التقنيين المخصص ولاختبار خريطة الطريق على النحو الذي فوض به الأطراف على النحو التالي:

(أ) في إطار المنتدى الإلكتروني المفتوح العضوية، عقد عشر مجموعات نقاش مخصصة وأربعة مؤتمرات إلكترونية إقليمية في الوقت الحقيقي (أفريقيا وآسيا والمحيط الهادي)؛¹¹

(ب) في إطار فريق الخبراء التقنيين المخصص: خمس جولات لمجموعات النقاش الإلكترونية ومؤتمران عن بعد لمكتب فريق الخبراء التقنيين واجتماعات مباشرة للفريق العامل الفرعي المعنى بخريطة الطريق ومكتب فريق الخبراء التقنيين المخصص.¹²

15. وكانت الأنشطة المشار إليها في الفقرة 14 تتم بالتناوب بين منتدى الخبراء الإلكتروني المفتوح وفريق الخبراء التقنيين المخصص وذلك لتكوين حلقة استرجاع معلومات لكل مشروع جديد من تسخ وثائق الإرشاد التي أعدتها مجموعات العمل الفرعية التابعة لفريق الخبراء التقنيين المخصص وللممكن من مشاركة عدد كبير من الخبراء في كافة جوانب العملية.

16. وعقد الاجتماع الثاني لفريق الخبراء التقنيين المخصص خلال الفترة من 20 إلى 24 أبريل/ نيسان 2010 في بلجيانيا، سلوفينيا. وحضر الاجتماع أربع عشر عضواً في الفريق من الأطراف فضلاً عن عضوين من غير الأطراف وأربعة من المنظمات.

17. ويتضمن المرفق الثاني بهذه المذكرة قائمة كاملة بالأنشطة التي نفذت في إطار المنتدى الإلكتروني وفريق الخبراء التقنيين المخصص.

⁸ تتوافر قائمة المشاركين على http://bch.cbd.int/onlineconferences/participants_ra.shtml.

⁹ ترصد النصوص الكاملة لمجموعات النقاش على http://bch.cbd.int/onlineconferences/archived_discussions_ra.shtml وتتوافر الوثائق والنصوص الكاملة للمؤتمر الإلكتروني في الوقت الحقيقي على http://bch.cbd.int/onlineconferences/realtime_ra.shtml.

¹⁰ الفقرة 18 من المرفق الثالث بالمقرر 10/8 لمؤتمر الأطراف

¹¹ تتوافر النصوص الكاملة لمجموعات النقاش على http://bch.cbd.int/onlineconferences/archived_discussions_ra.shtml وترد الوثائق والنصوص الكاملة للمؤتمرات الإلكترونية في الوقت الحقيقي على http://bch.cbd.int/onlineconferences/realtime_ra.shtml.

¹² عقدت اجتماعات الفريق العامل المعنى بخطة الطريق ومكتب فريق الخبراء التقنيين المخصص في لاهاي من 12 إلى 14 أكتوبر/ تشرين الأول 2009.

ألف- نتائج منتدى الخبراء الإلكتروني المفتوح العضوية المعنى بتقييم المخاطر وإدارة المخاطر

18. فيما يلي التوصيات الصادرة عن المنتدى الإلكتروني لفريق الخبراء التقنيين المخصص قبيل الاجتماع الأول:
- (أ) وضع إرشاد بشأن الجوانب النوعية التالية لتقييم المخاطر وإدارة المخاطر: (1) الأسماك والأشجار والكائنات الدقيقة والنباتات الطبية المحورة الحية؛ (2) الكائنات المحورة الحية المزودة بجينات أو سلالات مدعمة؛ (3) بيئات متلقية نوعية (4) الرصد بعد الإطلاق والتأثيرات الطويلة الأجل للكائنات المحورة الحية بعد إطلاقها في البيئة.
- (ب) خطة عمل لوضع مواد إرشادية بشأن الجوانب النوعية المرتبة بحسب الأولوية فضلا عن خريطة الطريق.
19. وبعد الاجتماع الأول لفريق الخبراء التقنيين المخصص، ساعدت المناقشات التي جرت في إطار منتدى الخبراء الإلكتروني المفتوح العضوية في وضع المشروع واختبار خريطة الطريق فضلا عن وضع الإرشاد الخاص بالجوانب النوعية لتقييم المخاطر التي حددها فريق الخبراء التقنيين المخصص باعتبارها من الأولويات (أي البعوض المحور الحي، والمحاصيل المحورة الحية التي تتحمل الإجهاد اللا أحيائي والكائنات المحورة الحية المزودة بجينات مدعمة.
20. وخلال العديد من جولات المناقشات، قدم الخبراء في المنتدى الإلكتروني مدخلات كبيرة في فريق الخبراء التقنيين المخصص بشأن محتويات خريطة الطريق والجوانب النوعية لتقييم المخاطر. ولدى اختبار خريطة الطريق، كانت غالبية الآراء تتسم بالإيجابية فيما يتعلق بفائدة العديد من الإرشادات التي قدمت وصلتها الوثيقة بالموضوع فيما يتعلق بالطرق المؤدية إلى تحسين الجوانب المواتية للمستخدمين إزاء خريطة الطريق.
21. وخلال الجولة الأخيرة لمجموعات النقاش المخصصة، دعي أعضاء المنتدى الإلكتروني إلى تقديم توصيات لاجتماع الأطراف للنظر خلال اجتماعه الخامس بشأن الطريق إلى الأمام فيما يتعلق بعمليات تقييم المخاطر وإدارة المخاطر. وأعرب المشاركون في المنتدى عن وجهة نظر بشأن خريطة الطريق والإرشاد المتعلق بالجوانب النوعية لتقييم المخاطر، وأشاروا إلى ضرورة تنقيح وتحديد هذه الوثائق بصورة منتظمة لضمان صلتها بالموضوع ومواكبة التطورات الجديدة.
22. كما أشار المشاركون في المنتدى الإلكتروني إلى الحاجة إلى وضع إرشاد إضافي بشأن الجوانب النوعية الأخرى لتقييم المخاطر. وأشار المنتدى إلى مواضيع تقييم المخاطر الواردة في وثائق المعلومات UNEP/CBD/BS/COP-MOP/5/INF/12 و UNEP/CBD/BS/COP-MOP/5/INF/13 باعتبارها نقطة البداية لوضع إرشاد آخر¹³. وعلاوة على ذلك، أوصى المشاركون أيضا بالنظر في المواضيع التالية: (1) وضع سيناريوهات للمخاطر؛ (2) استراتيجيات لإدارة المخاطر بما في ذلك رصد تأثيرات الكائنات المحورة الحية بعد إطلاقها في البيئة؛ (3) تحليل الشوك والتقلبات؛ (4) "قائمة مراجعة تحتوي على العناصر الحرجة في عملية تقييم المخاطر؛ (5) الطريقة الأفضل لربط عملية تقييم المخاطر في إطار البروتوكول بالأحكام والمقررات في إطار اتفاقية التنوع البيولوجي.
23. وأوصى كذلك، خلال مناقشات المنتدى الإلكتروني بمواصلة التشاور، أثناء وضع الإرشاد الجديد، فيما بين الأطراف، وضرورة مراعاة الإرشاد الحالي الذي أعدته الهيئات الدولية الأخرى (منظمة التعاون والتنمية في الميدان الاقتصادي والاتفاقية الدولية لوقاية النباتات.
24. وفيما يتعلق بآلية لمعالجة عملية وضع الإرشاد الآخر، أوصى عدد كبير من الخبراء بإجراء مناقشات إلكترونية بواسطة فريق الخبراء التقنيين المخصص وتبادل المعلومات من خلال غرفة تبادل معلومات السلامة الأحيائية أو توليفة منهما. وتتضمن الأمثلة الأخرى على الآليات الخاصة بمعالجة وضع الإرشاد إجراء مشاورات فيما بين الخبراء وتجميع لخبراء المعلومات لتنفيذ تدريب المتابعة بعد وضع الإرشاد.
25. وقد جمعت وجهات النظر والتوصيات التي صدرت في إطار منتدى الخبراء الإلكتروني المفتوح العضوية وأتيحت بوصفها وثائق معلومات للنظر من جانب الأطراف (UNEP/CBD/BS/COP-MOP/5/INF/12 و 14)¹⁴.

¹³ يتوافر على <http://bch.cbd.int/protocol/meetings/documents.shtml?eventid=3018>

¹⁴ تتوافر وثائق المعلومات UNEP/CBD/BS/COP-MOP/5/INF/12 و UNEP/CBD/BS/COP-MOP/5/INF/14 على <http://bch.cbd.int/protocol/meetings/documents.shtml?eventid=3018>

باء- نتائج فريق الخبراء التقنيين المخصص المعنى بتقييم المخاطر وإدارة المخاطر

26. تتمثل النتائج الرئيسية للاجتماع الأول لفريق الخبراء التقنيين المخصص المعنى بتقييم المخاطر وإدارة المخاطر فيما يلي: (1) مشروع خريطة الطريق؛ (2) تحديد وترتيب أولويات المسائل النوعية الأخرى لتقييم المخاطر (أي البعوض المحور الحي والمحاصيل المحورة الحية مع تحمل الإجهاد اللا أحيائي والكائنات المحورة الحية المزودة بجينات مدعمة) لأغراض وضع الإرشاد؛ (3) إنشاء أربعة أفرقة عمل فرعية للتركيز على كل مسألة من المسائل المحددة؛ (4) وضع خطة عمل مكونة من موجز للمصطلحات والإجراءات الخاصة بوضع إرشاد قبل الاجتماع الثاني لفريق الخبراء التقنيين المخصص.

27. وخلال فترة ما بين الدورات، واصلت أفرقة العمل الفرعية لفريق الخبراء التقنيين المخصص، من خلال مشاورات مع فريق الخبراء الإلكتروني المفتوح العضوية، وضع مشروع الوثائق الخاصة بالإرشاد المتعلق بالمسائل النوعية الأربعة لتقييم المخاطر ومشروع خريطة الطريق المجرية بشأن تقييم مخاطر الكائنات المحورة الحية.

28. وكانت النتائج الرئيسية لفريق الخبراء التقنيين المخصص خلال اجتماعه الثاني ما يلي:

(أ) الانتهاء من الوثيقة المعنونة "إرشاد بشأن تقييم مخاطر الكائنات المحورة الحية" وتقسيمها إلى قسمين بعنوان "القسم الأول: خريطة طريق لتقييم مخاطر الكائنات المحورة الحية" والقسم الثاني "الأنواع المحددة للكائنات والسلالات المحورة الحية" (أي المحاصيل المحورة الحية مع تحمل الإجهاد اللا أحيائي، والبعوض المحور الحي والكائنات المحورة الحية المزودة بجينات أو سلالات مدعمة). وترد هذه الوثيقة رفق هذه المذكرة في الملحق الثالث، وسوف تتوافر أيضا من خلال غرفة تبادل معلومات السلامة الأحيائية؛¹⁵

(ب) توصيات للأمانة بشأن كيفية دمج وتحديث وثيقة الإرشاد التي أعدها فريق الخبراء التقنيين المخصص والأدوات الخاصة باسترجاع مواد المعلومات الأساسية المتوافرة في مركز موارد معلومات السلامة الأحيائية في غرفة تبادل معلومات السلامة الأحيائية؛

(ج) تقييم لخطة العمل التي وضعت خلال الاجتماع الأول.

29. ووضع فريق الخبراء التقنيين المخصص أيضا توصيات للأطراف خلال اجتماعها الخامس لمواصلة إعداد إرشاد بشأن المواضيع الإضافية لتقييم المخاطر ولأسيما تلك المسائل النوعية لتقييم المخاطر التي جرى تحديدها وترتيب أولوياتها خلال المنتدى الإلكتروني المفتوح العضوية، والاجتماع الأول لفريق الخبراء التقنيين المخصص.

30. ويتوافر تقرير الاجتماع الأول والتقرير الأخير لفريق الخبراء التقنيين المخصص في وثائق معلومات للنظر من جانب الأطراف.¹⁶

31. وترفق المجموعة الكاملة للتوصيات المقدمة من فريق الخبراء التقنيين المخصص إلى الاجتماع الخامس للأطراف بهذه المذكرة في الملحق الرابع.

ثالثا - بناء القدرات في مجال تقييم المخاطر

32. استجابة لطلب الأطراف بشأن بناء القدرات في مجال تقييم المخاطر، قامت الأمانة بتنسيق عملية متعددة أصحاب المصلحة لتنظيم التدريب بالتعاون مع منظمات الأمم المتحدة (اتفاقية أروهوس للجنة الأمم المتحدة الاقتصادية لأوروبا والاتفاقية الدولية لوقاية النباتات لمنظمة الأغذية والزراعة، وبرنامج الأمم المتحدة للبيئة) وغيرها من المنظمات الدولية (الائتلاف العالمي للصناعة وشبكة العالم الثالث) والقطاع الأكاديمي (جامعة كانتربوري وجامعة مينيسوتا).

33. وقد جرى إعداد التدريب بطريقة تدريجية. فقد أعدت الأمانة أولا مخططا للتدريب ودعت المتعاونين إلى توفير المدخلات والتعليقات. وبعد ذلك وعلى أساس مختلف المعلومات المسترجعة، أعدت الأمانة مشروع دليل تدريبي ودعت المتعاونين إلى إجراء استعراض نظير. وقامت الأمانة بعد ذلك بتنقيح مشروع الدليل في ضوء المعلومات المسترجعة والتعليقات المقدمة خلال عملية الاستعراض النظير.

¹⁵ تتوافر على http://bch.cbd.int/onlineconferences/forum_RA.shtml

¹⁶ تتوافر الوثيقتان UNEP/CBD/BS/COP-MOP/5/INF/13 و UNEP/CBD/BS/COP-MOP/5/INF/15 على <http://bch.cbd.int/protocol/meetings/documents.shtml?eventid=3018>

34. وفي حين استخدمت الأمانة أحكام اتفاقية قرطاجنة للسلامة الأحيائية ولاسيما الملحق الثالث فيه كأساس لصيانة واستعراض دليل التدريب الناشئ، حاولت أيضا إدراج خبرات وممارسات جارية من عدد من الأطر التنظيمية الوطنية والمنظمات الدولية بطريقة شاملة.

35. وقد أسفرت هذه العملية عن مشروع لدليل التدريب بعنوان "تقييم مخاطر الكائنات المحورة الحية" يتألف من أربع وحدات هي (1) عرض عام للسلامة الأحيائية وبروتوكول قرطاجنة للسلامة الأحيائية (2) عمل تحضيرى- فهم السياق الذي تجري فيه عملية تقييم المخاطر (3) إجراء تقييم المخاطر (4) إعداد تقرير تقييم المخاطر.

36. ويتوافر دليل التدريب كوثيقة معلومات ومن خلال غرفة تبادل معلومات السلامة الأحيائية للنظر من جانب الأطراف.¹⁷

37. ولمواصل معالجة طلب الأطراف تنظيم أنشطة لبناء القدرات بغرض تمكين البلدان من تبادل الخبرات واكتساب المعارف المباشرة بشأن إعداد وتقييم تقارير تقييم المخاطر وفقا لأحكام البروتوكول، استخدم دليل التدريب المشار إليه أعلاه خلال الأنشطة التالية:

(أ) حلقة العمل دون الإقليمية للمحيط الهادي بشأن بناء القدرات وتبادل الخبرات المتعلقة بتقييم المخاطر التي عقدت في نادي، فيجي خلال الفترة من 4 إلى 7 يوليو/ تموز 2010؛

(ب) حلقة العمل دون الإقليمية في آسيا بشأن تقييم مخاطر الكائنات المحورة الحية التي عقدت في سيام ريب، كمبوديا من 12 إلى 16 يوليو/ تموز 2010.

38. وقد حضر حلقة العمل دون الإقليمية للمحيط الهادي اثني عشر مشاركا من ستة أطراف في البروتوكول (فيجي وكيريباتي ونيوي وساموا وجزر سليمان وتونغا) وبلدان من غير الأطراف (جزر كوك وفانواتو) ومنظمة واحدة (جامعة كانتربروري، نيوزيلندا) وحضر الدورة التدريبية لآسيا ثلاث وعشرون مشاركا من خمس عشر طرفا في البروتوكول (بهوتان وكمبوديا، والهند، وأندونيسيا، وجمهورية إيران الإسلامية، وجمهورية لاو الديمقراطية الشعبية وماليزيا، ومنغوليا، وميانمار، وباكستان والجمهورية العربية السورية، وتايلند، وتوركمنستان، وفيتنام واليمن)، ومنظمة غير حكومية (شبكة العالم الثالث) وبرنامج الأمم المتحدة للبيئة. كما شارك في الدورة التدريبية لآسيا محاضر من هولندا.

39. ودعي المشاركون إلى الإجابة على استبيان لتقييم حلقة عمل المحيط الهادي والدورة التدريبية لآسيا. وأشارت نتائج الاستبيان إلى اتفاق عام على أن هذين النشاطين (1) قدما تدريبا مباشرا في مجال إعداد وتقييم تقارير تقييم المخاطر وفقا لمواد البروتوكول والملحق الثالث به (2) ساعدا على تنمية المهارات بشأن كيفية استخدام وتفسير المعلومات الحالية فضلا عن تحديد ومعالجة الثغرات في المعلومات (3) ساعدا في فهم كيفية الثغرات وضع معلومات خط الأساس ذات الصلة بتقييم المخاطر.

40. وأوضحت نتائج الاستبيان أيضا أن غالبية المشاركين يتفقون على أن دليل التدريب الذي أعدته الأمانة بالتعاون مع هيئات الأمم المتحدة الأخرى والمنظمات ذات الصلة (1) أداة مفيدة للتدريب على تقييم المخاطر (2) سهل الفهم بطريقة تدريجية (3) يتألف من عرض عام كاف لعملية تقييم المخاطر و(4) مفيد لطائفة عريضة من المستخدمين.

41. ورأى المشاركون، لدى تقديم المزيد من المعلومات المسترجعة، أن دليل التدريب يمثل أداة تعليمية ممتازة توفر تقدما حسن التنظيم وشاملا لعملية تقييم المخاطر ومفيد للأطراف فضلا عن البلدان الأخرى والمنظمات ذات الصلة. وأشار المشاركون إلى أنه يتعين، لتحسين فائدة دليل التدريب مايلي:

(أ) زيادة تحسين الدليل من خلال جملة أمور من بينها إضافة مسرد للمصطلحات، وقائمة بالمترادفات ومخططات التدفق والأشكال البيانية وأمثلة على الكائنات المحورة الحية غير المحصولية وغير ذلك؛

(ب) إدراج عناصر من "الإرشاد الخاص بتقييم مخاطر الكائنات المحورة الحية" الذي وضعه فريق الخبراء التقنيين المخصص من خريطة الطريق أساسا (مثل مخطط التدفق) ومن الإرشاد بشأن الأنواع المحددة من الكائنات والسلالات المحورة الحية (أي تقييم مخاطر البعوض المحور الحي والكائنات المحورة الحية المزودة بجينات أو سلالات مدعمة والمحاصيل المحورة الحية التي تتحمل الإجهاد اللا أحيائي؛

(ج) تقديم الدليل من خلال أداة تعلم صديقة للمستخدمين (مثل البرمجيات التفاعلية)؛

¹⁷ يتوافر دليل التدريب في وثيقة المعلومات at UNEP/CBD/BS/COP-MOP/5/INF/22 على

<http://bch.cbd.int/protocol/meetings/documents.shtml?eventid=3018> http://bch.cbd.int/protocol/cpb_art15/training

(د) إصداره في جميع لغات الأمم المتحدة.

42. واتفق المشاركون في حلقة عمل المحيط الهادي والدورة التدريبية لآسيا على أنه يمكن للأطراف خلال اجتماعها الخامس النظر في العناصر والأنشطة التالية:

بناء القدرات في مجال تقييم المخاطر

(أ) المزيد من الدورات التدريبية بشأن تقييم المخاطر على المستوى الوطني أو للمناطق الجغرافية الصغيرة (مثل نحو 5 إلى 7 بلدان) حيث تتماثل البيئة المتلقية مما يتيح مشاركة الفريق الأساسي من الخبراء القطريين في كل بلد؛

(ب) تدريب متابعة متقدم في مجال المخاطر بالتركيز، على سبيل المثال، على مختلف أنواع الاستخدامات المتوخاة (أي الإدخال في البيئة والكائنات الحية لأغراض الاستخدام المباشر كأغذية وأعلاف أو للتصنيع) ومختلف أنواع الكائنات المحورة الحية؛

(ج) دورات تدريبية مخصصة بشأن (1) إعداد تقارير تقييم المخاطر والتوصيات المتصلة بها (2) استخلاص البيانات ذات الصلة من الإخطارات (3) تقييم نوعية البيانات المقدمة في الطلب و(4) وضع معلومات مفصلة لخط الأساس؛

(د) تدريب المدربين الذين يمكنهم الاضطلاع بعمليات بناء القدرات على المستوى الوطني.

إرشاد بشأن تقييم المخاطر:

(هـ) نشر وتوزيع الوثيقة "إرشاد بشأن تقييم مخاطر الكائنات المحورة الحية بما في ذلك النسخة الإلكترونية على غرفة تبادل معلومات السلامة الأحيائية بجميع لغات الأمم المتحدة؛

(و) وضع إرشاد آخر بشأن تقييم المخاطر على النحو الذي أوصى به فريق الخبراء التقنيين المخصص؛

بناء القدرات بشأن السلامة الأحيائية

(ز) المزيد من التدريب الإقليمي بشأن تحديد هوية الكائنات المحورة الحية؛

(ح) تدريب صانعي القرار على تفسير التوصيات الخاصة بتقييم المخاطر وبشأن تنفيذ استراتيجيات إدارة المخاطر.

43. وتتوافر تقارير أنشطة بناء القدرات هذه في شكل وثائق ومعلومات للنظر من جانب الأطراف (UNEP/CBD/BS/COP-MOP/5/INF/16 و17).¹⁸

رابعاً - التعاون في تحديد هوية الكائنات المحورة الحية أو السلالات النوعية التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر

44. دعا الأمين التنفيذي، في أخطار، الأطراف والحكومات الأخرى والمنظمات ذات الصلة إلى تقديم معلومات سليمة من الناحية العملية عن تحديد هوية الكائنات المحورة الحية أو السلالات النوعية التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر.¹⁹

45. أشير في بعض التقديمات التي تلقتها الأمانة إلى الكائنات المحورة الحية أو السلالات النوعية التي قد يكون لها تأثيرات معاكسة مثل القطن والأسماك والذرة والأشجار والفيروسات المحورة الحية فضلاً عن الكائنات المحورة الحية لإنتاج المستحضرات الصيدلانية مع جينات أو سلالات مدعمة ومقاومة الحشرات وتحمل الإجهاد اللا أحيائي والمبيدات. وأشارت بعض التقديمات من ناحية أخرى إلى أنه لا تتوافر أي قرائن تعتمد على العلم تشير إلى التأثيرات المعاكسة المحتملة للكائنات المحورة الحية المتداولة في التجارة حتى الآن.

¹⁸ تتوافر وثيقتنا المعلومات UNEP/CBD/BS/COP-MOP/5/INF/16 وUNEP/CBD/BS/COP-MOP/5/INF/17 على الموقع

<http://bch.cbd.int/protocol/meetings/documents.shtml?eventid=3018>

¹⁹ الاخطار (2009-056) SCBD/BS/MPDM/jh/67587 متوافر على <http://bch.cbd.int/protocol/notifications/>.

46. وبناء على التقديرات المشار إليها أعلاه، أعدت الأمانة وثيقة بعنوان "تجميع للتقديرات عن تحديد هوية الكائنات المحورة الحية أو السلالات النوعية التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضا المخاطر على صحة البشر" للنظر من جانب فريق الخبراء التقنيين المخصص والأطراف.²⁰

47. ويعد مداولات بشأن هذه المسألة، حدد فريق الخبراء التقنيين المخصص الطرائق التالية للتعاون (1) تبادل المعلومات عن طريق غرفة تبادل معلومات السلامة الأحيائية (2) حلقات العمل (3) فريق خبراء تقنيين مخصص و(4) التعاون في اختبار الكائنات المحورة الحية.

48. كما وافق عدد من أعضاء فريق الخبراء التقنيين المخصص على إمكانية إنشاء عملية تدريجية لهذا الغرض حيث يعقب المرحلة الأولى لتجميع المعلومات مرحلة ثانية لتحليل المعلومات.

49. وقدم فريق الخبراء التقنيين المخصص توصيات نوعية أخرى فيما يتعلق بهذه المسألة على النحو الوارد في الفقرات (و) و(ز) و(4) من الملحق أدناه.

خامسا- تحديد هوية الكائنات المحورة الحية التي ليس لها تأثيرات معاكسة على الأراجح على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضا المخاطر على صحة البشر

50. تنص الفقرة 4 من المادة 7 من البروتوكول على أن إجراء "الموافقة المسبقة عن علم لن يسري على حركة الكائنات المحورة الحية المتعمدة عبر الحدود المحددة في مقرر لمؤتمر الأطراف العاملة كاجتماع للأطراف في البروتوكول بأنها قد لا تنطوي على تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضا المخاطر على صحة البشر".

51. يمكن للأطراف في الاجتماع الخامس لدى مداولاتها بشأن الطرائق التي قد تمكن من تحديد هوية الكائنات المحورة الحية التي قد لا تنطوي على تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضا المخاطر التي تتعرض لها صحة البشر، أن تأخذ في الاعتبار، ضمن جملة أمور، التقديرات التالية من الأطراف من خلال غرفة تبادل معلومات السلامة الأحيائية في إطار الإجراء المبسط (المادة 13) الذي جرى فيه إعفاء الواردات من الكائنات المحورة الحية من إجراء الموافقة المسبقة عن علم²¹.

52. واعتبارا من 10 يونيو/ حزيران 2010، قدمت الكائنات المحورة الحية التالية إلى غرفة تبادل معلومات السلامة الأحيائية بمقتضى الإجراء المبسط.

الكائنات المحورة الحية التي طبق عليها الإجراء المبسط	البلد	سجل غرفة تبادل المعلومات
قطن بولجارد	كولومبيا	8151
القطن جاهز جينات المقاومة	كولومبيا	8155
قطن بولجارد (MON-15985-7)	جنوب أفريقيا	5666
قطن بولجارد (MON-00531-6)	جنوب أفريقيا	5679
ذرة البيلد جارد (MON-00810-6)	جنوب أفريقيا	5712
ذرة البيلد جارد (SYN-BT011-1)	جنوب أفريقيا	5715
ذرة البيلد جارد (MON-00603-6)	جنوب أفريقيا	8164
فول الصويا جاهز جينات المقاومة (MON-04032-6)	جنوب أفريقيا	8167
القطن جاهز جينات المقاومة (MON-01445-2)	جنوب أفريقيا	8170

²⁰ تتوافر الوثيقة UNEP/CBD/BS/COP-MOP/5/INF/11 كوثيقة معلومات على <http://bch.cbd.int/protocol/meetings/documents.shtml?eventid=3018>.

²¹ المادة 13 الفقرة 1 (ب).

سجل غرفة تبادل المعلومات	البلد	الكائنات المحورة الحية التي طبق عليها الإجراء المبسط
40513	جنوب أفريقيا	الذرة البيلد جارد جاهز جينات المقاومة (MON-00603-6 x MON-00810-6)
40514	جنوب أفريقيا	قطن الفليكس جاهز جينات المقاومة (MON-88913-8)
40516	جنوب أفريقيا	قطن البيلد جارد جاهز جينات المقاومة (MON-00531-6 x MON-01445-2)

سادسا- الاستنتاجات وعناصر مشروع مقرر

ألف- الإرشاد الآخر بشأن الجوانب النوعية لتقييم المخاطر

53. نفذت بنجاح المهام التي أوكلتها الأطراف في اختصاصات المنتدى الإلكتروني وفريق الخبراء التقنيين المخصص بشأن وضع إرشاد آخر يتعلق بتقييم المخاطر من خلال عملية تضمنت كلا من المداولات بالوسائل الإلكترونية والمباشرة.

54. وقد تداولت مجموعة كبيرة من الخبراء الكثرونيا عبر مجموعات النقاش المخصصة والمؤتمرات في الوقت الحقيقي، وقدمت توصيات إلى مجموعة أصغر هي فريق الخبراء التقنيين المخصصة اجتمعت بصورة مباشرة. وقد أتاحت هذه العملية لعدد كبير من الخبراء في مختلف الميادين العلمية والتقنية ذات الصلة بتقييم المخاطر توفير مدخلات في عملية وضع المواد الإرشادية بطريقة تحقق مردودية تكاليفها في حدود الموارد المالية المحدودة المتاحة.

55. وقد أسفرت هذه العملية عن وثيقة بعنوان "إرشاد بشأن تقييم مخاطر الكائنات المحورة الحية". وقد أوصى فريق الخبراء التقنيين المخصص والمنتدى الإلكتروني بضرورة معاملة وثيقة الإرشاد على النحو التالي: (1) الإصدار والتوزيع بما في ذلك النسخة الإلكترونية في غرفة تبادل معلومات السلامة الأحيائية بجميع لغات الأمم المتحدة (2) مواصلة تجربتها وذلك مثلا خلال حلقات العمل الإقليمية بما في ذلك التعاون مع المبادرات العاملة لبناء القدرات والتدريب حسب مقتضى الحال (3) تنقيحها في غضون عامين والحاجة إلى تحديث قائمة مواد المعلومات الأساسية في غضون عام.

56. وفي حين أحرز تقدم كبير في معالجة الحاجة إلى إرشاد بشأن تقييم المخاطر من خلال إعداد الوثيقة المشار إليها أعلاه، كان من رأي الكثير من أعضاء فريق الخبراء التقنيين المخصص والمنتدى الإلكتروني أن الأمر مازال يحتاج إلى مواصلة عملية الجمع بين منتدى الكثروني وفريق خبراء تقنيين مخصص.

57. واستنادا إلى المعلومات الواردة أعلاه ومراعاة ضمن جملة أمور لتوصيات المنتدى الإلكتروني وفريق الخبراء التقنيين المخصص، قد يرغب مؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول فيما يلي:

(أ) أن يؤيد مواصلة عمل كل من منتدى الخبراء الإلكتروني المفتوح العضوية وفريق الخبراء التقنيين المخصص المعني بتقييم المخاطر وإدارة المخاطر والموافقة على ذلك من أجل: (1) وضع إرشاد إضافي بشأن الأنواع المحددة من الكائنات والسلالات المحورة الحية مع مراعاة ضمن جملة أمور المواضيع المدرجة في الملحق الخامس أدناه و(2) تنقيح نص "الإرشاد بشأن تقييم مخاطر الكائنات المحورة الحية" وذلك مثلا في ضوء اختبار الإرشاد خلال أنشطة بناء القدرات وتحديث قوائمه الخاصة بمواد المعلومات الأساسية؛

(ب) أن يطلب إلى الأمين التنفيذي (1) إصدار وتوزيع وثيقة الإرشاد بشأن تقييم مخاطر الكائنات المحورة الحية: بجميع لغات الأمم المتحدة بما في ذلك النسخة الإلكترونية في غرفة تبادل معلومات السلامة الأحيائية (2) إختبار وثيقة الإرشاد خلال حلقات العمل الإقليمية بما في ذلك التعاون مع المبادرات الجارية بشأن بناء القدرات والتدريب حسب مقتضى الحال (3) تنقيح النموذج الموحد لتقديم السجلات إلى مركز موارد معلومات السلامة الأحيائية في غرفة تبادل معلومات السلامة الأحيائية سعيا إلى ربط سجلات المركز بشأن تقييم المخاطر بالأقسام النوعية في وثيقة الإرشاد؛

(ج) أن يواصل المناقشات في إطار منتدى الخبراء الإلكتروني المفتوح العضوية بشأن تقييم المخاطر وإدارة المخاطر، وأن يطلب إلى الأمين التنفيذي إصدار الدعوة إلى خبراء آخرين؛

(د) أن ينشئ فريق خبراء تقنيين مخصص معني بتقييم المخاطر وإدارة المخاطر وأن يطلب إلى الأمين التنفيذي تطبيق نفس الإجراءات مع تعديلها حسب الظروف لاختيار الخبراء مثلما كان الحال في العملية السابقة.

باء- بناء القدرات على تقييم المخاطر

58. فيما يتعلق ببناء القدرات، وضع دليل تدريب بالتعاون مع بعض منظمات الأمم المتحدة والمنظمات الدولية المعنية. وقد استخدم الدليل كأساس لأنشطة بناء القدرات التي جرت في الإقليميين الفرعيين للمحيط الهادي وآسيا. وقد المشاركون في حلقة العمل والدورة التدريبية العديد من التوصيات بشأن تحسين دليل التدريب فيما يتعلق بفوائده وخطوطه المواتية للمستخدمين. وعلاوة على ذلك، أوصى المشاركون بتطوير الدليل بوصفه مواد تدريب تفاعلية (أي على قرص مغمط) وترجمته إلى جميع لغات الأمم المتحدة.

59. واستناداً إلى المعلومات الواردة أعلاه ومراعاة لجملة أمور من بينها توصيات المشاركين في أنشطة بناء القدرات، قد يرغب مؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول:

(أ) أن يطلب إلى الأمين التنفيذي أن يعقد في أقرب وقت ممكن ورهنا بتوافر الأموال، المزيد من الدورات التدريبية الإقليمية ودون الإقليمية لتمكين البلدان من اكتساب خبرات مباشرة في إعداد وتقييم تقارير تقييم المخاطر وفقاً لمواد البروتوكول والملحق الثالث؛

(ب) أن يطلب كذلك إلى الأمين التنفيذي أن يحسن، بالتعاون مع منظمات الأمم المتحدة والمنظمات الأخرى ذات الصلة من فائدة دليل التدريب بشأن "تقييم مخاطر الكائنات المحورة الحية" من خلال (1) التنقيح المنتظم للدليل في ضوء التوصيات التي قدمت خلال أنشطة بناء القدرات الإقليمية ودون الإقليمية (2) تطوير الدليل ليصبح أداة تفاعلية للتعلم مثل القرص المغمط وإتاحتها من خلال غرفة تبادل معلومات السلامة الأحيائية (3) إصدار وتوزيع الدليل على الأطراف والحكومات الأخرى والمنظمات ذات الصلة.

جيم- تحديد هوية الكائنات المحورة الحية أو السلالات النوعية التي (1) قد تنطوي أو (2) لا تنطوي على تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة المخاطر على صحة البشر

60. أعربت الأطراف والحكومات الأخرى والمنظمات المعنية عن وجهات نظر متباينة فيما يتعلق بتحديد هوية الكائنات المحورة الحية أو السلالات النوعية التي قد يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضاً المخاطر على صحة البشر. وحدد فريق الخبراء التقنيين المخصص الطرائق التالية لمعالجة المسألة (1) تبادل المزيد من المعلومات من خلال غرفة تبادل معلومات السلامة الأحيائية (2) حلقات العمل (3) فريق خبراء تقنيين مخصص و(4) التعاون في معالجة التأثيرات المعاكسة المحتملة للكائنات المحورة الحية. ويمكن بدء هذه العملية بطريقة تدريجية من خلال المرحلة الأولى لجمع المعلومات يعقبها تحليل للمعلومات.

61. وفيما يتعلق بتحديد هوية الكائنات المحورة الحية التي قد لا تنطوي على الأرجح على تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضاً المخاطر على صحة البشر، قد تأخذ الأطراف علماً، ضمن جملة أمور، بالمقررات التي اتخذت في إطار الإجراء المبسط بشأن الواردات من الكائنات المحورة الحية المعفاة من إجراء الموافقة المسبقة عن علم. وتقديمها لغرفة تبادل معلومات السلامة الأحيائية.

62. واستناداً إلى المعلومات الواردة أعلاه ومع الأخذ في الاعتبار، ضمن جملة أمور، وجهات النظر التي أبدتها الأطراف والحكومات الأخرى والمنظمات المعنية، وتوصيات المنتدى الإلكتروني المفتوح العضوية وفريق الخبراء التقنيين المخصص، قد يرغب مؤتمر الأطراف العامل كاجتماع للأطراف في البروتوكول في إنشاء آلية أو أكثر بما في ذلك على سبيل المثال تبادل المعلومات وحلقات العمل وفريق خبراء بهدف تمكين الأطراف من اتخاذ المقررات بشأن تحديد هوية الكائنات المحورة الحية أو السلالات النوعية التي (1) قد يكون (2) أو لا يكون لها تأثيرات معاكسة على صون التنوع البيولوجي واستخدامه المستدام مع مراعاة أيضاً المخاطر على صحة البشر.

*Annex I***LIST OF AHTEG MEMBERS****PARTIES****Austria**

1. Dr. Helmut Gaugitsch
Head of Unit
Landuse & Biosafety
Federal Environment Agency
Spittelauer Lände 5
Vienna A-1090, Austria
Tel.: +43 1 31 304 3133
Fax: +43 1 31 304 3700
E-Mail: helmut.gaugitsch@umweltbundesamt.at
Web: <http://www.umweltbundesamt.at>

Belize

2. Dr. Michael DeShield
Director
Food Safety Services
Belize Agricultural Health Authority
Central Investigation Laboratory
P.O. Box 181
Belize City, Belize
Tel.: +501 224 4794
Fax: +501 224 5230
E-Mail: foodsafety@btl.net, deshield@btl.net

Brazil

3. Dr. Eliana Maria Gouveia Fontes
Senior Scientist
Biological Control Unit /Ecology,
Semiocemicals & Biosafety Laboratory
EMBRAPA-Cenargen
C.P. 02372
Brasilia, DF 71.510-230, Brazil
Tel.: +55 61 448 4793
Fax: +55 61 3448 4672
E-Mail: efontes@cnpq.br,
efontes551@gmail.com

China

4. Mr. Wei Wei
Associate Professor
Institute of Botany
Chinese Academy of Sciences
20 Nanxincun, Xiangshan, Beijing 100093, China
Tel.: +86 10 6283 6275
Fax: +86 10 8259 6146
E-Mail: weiwei@ibcas.ac.cn

Croatia

5. Ms. Jelena Zafran Novak
Expert
Laboratory for GMO Detection
Croatian National Institute of Public Health
Rockefellerova 7
Zagreb 10000, Croatia
Tel.: +385 1 4863207
Fax: +385 91 8996420
E-Mail: j.zafran-novak@hzjz.hr

Cuba

6. Prof. Leticia Pastor Chirino
Head
Department of Authorizations
National Centre for Biological Safety
Edif. 70c, apto 3. Zona 6 Alamar
Habana del este Ciudad Habana
Cuba
Tel.: +537 765 1202
Fax: +537 202 3255
E-Mail: leticia.ch@orasen.co.cu,
lpch06@yahoo.es

Egypt

7. Dr. Ossama Abdel-Kawy
Scientific Advisor
Egyptian Environmental Affairs Agency
30 Maadi Zerae Road, 7th Floor
Maadi, Cairo 12551, Egypt
Tel.: +20 11 561 456
E-Mail: elkawyo@gmail.com,
abdkawy@yahoo.com
Web: <http://eg.biosafetyclearinghouse.net>

Germany

8. Dr. Beatrix Tappeser
Head of Division
Biosafety. GMO Regulation
Federal Agency for Nature Conservation
Konstantinstr. 110, Bonn D-53179, Germany
Tel.: +49 228 8491 1860
Fax: +49 227 8491 1869
E-Mail: TappeserB@bfn.de
Web: www.bfn.de

Japan

9. Prof. Kazuo Watanabe
 Professor, Plant Genetic Diversity,
 Biosafety and Bioethics
 Gene Research Center, University of Tsukuba
 Ministry of Education, Culture, Sports, Science
 and Technology
 1-1-1 Tennoudai, Tsukuba, Ibaraki 305-8572,
 Japan
 Tel.: +81 29 853 4663
 Fax: +81 29 853 7723
 E-Mail: nabechan@gene.tsukuba.ac.jp

Malaysia

10. Dr. Chan Kok Gan
 Senior Lecturer, Genetics & Molecular Biology
 Faculty of Science
 University of Malaya
 Kuala Lumpur 50603, Malaysia
 Tel.: +603 7967 5162
 Fax: +603 7967 4509
 E-Mail: kokgan@um.edu.my
11. Dr. Vilasini Pillai
 Scientist in Residence
 Office of the Science Advisor
 Ministry of Science, Technology and
 Innovation
 Level 1-7, Block C5, Parcel C
 Federal Government Administrative Centre
 Putrajaya 62662, Malaysia
 Tel.: +6 03 8885 8707
 Fax: +6 03 8888 3801
 E-Mail: vilasini@mosti.gov.my
 Web: www.moste.gov.my

Mexico

12. Dra. Sol Ortiz Garcia
 Technical Director
 Comisión Intersecretarial de Bioseguridad
 de los Organismos Genéticamente
 Modificados
 San Borja 938, esquina Heriberto Frías, Colonia
 del Valle, delegación Benito Juárez
 México D.F. Distrito Federal – 03100, Mexico
 Tel.: +52 55 5575 7618 ext 22
 Fax: +52 55 5575 7618 ext 30
 E-Mail: sortiz@conacyt.mx

Netherlands

13. Dr. Hans Bergmans
 Senior Scientist
 SEC/GMO Office
 National Institute of Public Health and
 Environment
 Antonie van Leeuwenhoeklaan 9, PO Box 1
 Bilthoven 3720 BA, Netherlands
 Tel.: +31 30 274 4195, +6 20 737792
 Fax: +31 30 2744401
 E-Mail: hans.bergmans@rivm.nl

Niger

14. Mr. Gado Zaki Mahaman
 Direction Générale de l'Environnement et
 des Eaux et Forêts
 P.O. Box 721, Niamey, Niger
 Tel.: + 22796110415, +22720723755
 Fax: +227 20723763
 E-Mail: mahamane_gado@yahoo.fr

Nigeria

15. Mr. Rufus Ebegba
 Chief Environmental Scientist
 Federal Ministry of Environment
 Independence Way (South)
 Central Area, P.M.B. 468
 Garki-Abuja, Nigeria
 Tel.: +234 803 314 7778
 Fax: +234 9 523 4119
 E-Mail: rebegba@hotmail.com
16. Ms. Hajara Yusuf Sadiq
 Scientific Officer
 Environmental Biotech/Biosafety Unit
 National Biotechnology Development Agency
 16, Dunukofia Str. Area 11
 P.M.B. 5118, Wuse Zone 5
 Garki - Abuja FCT, Nigeria
 Tel.: +2348055179400, +2348066042543
 Fax: +234093145473
 E-Mail: haj4sadiq@yahoo.com

Norway

17. Dr. David Quist
Senior Scientist
Genome Ecology Section
GenØk – Centre for Biosafety
Science Park, PO 6418
Tromsø N-9294, Norway
Tel.: +47 77 646294
Fax: +47 77 646100
E-Mail: david.quist@uit.no

Republic of Moldova

18. Dr. Angela Lozan
Head of the Biosafety Office
Ministry of Environment
Str. Cosmonautilor 9, Bir 526
Chisinau, Republic of Moldova
Tel.: +373 22 22 68 74
Fax: +373 22 22 68 74
E-Mail: angelalozan@yahoo.com

Slovenia

19. Dr. Branka Javornik
National Expert - Professor of Genetics &
Biotechnology
Department of Agronomy, Biotechnical Faculty
University of Ljubljana
Jamnibarjeva 101
Ljubljana 1000, Slovenia
Tel.: +3861 423 1161
Fax: +3861 423 1088
E-Mail: branka.javornik@bf.uni-lj.si

NON-PARTIES

Australia

20. Dr. Paul Keese
Science Advisor
Office of the Gene Technology
Regulator
Department of Health and Ageing
MDP 54, GPO Box 9848
Canberra ACT 2601, Australia
Tel.: +61 2 6271 4254
Fax: +61 2 6271 4202
E-Mail: paul.keese@health.gov.au

Canada

21. Mr. Philip Macdonald
National Manager
Plant and Biotechnology Risk Assessment Unit
Canadian Food Inspection Agency
1400 Merivale Rd
Ottawa, ON K1A 0Y9, Canada
Tel.: +613 773 5288
Fax: +613 773 5391
E-Mail: philip.macdonald@inspection.gc.ca

United States of America

22. Mr. David Heron
Assistant Director Policy Coordination,
Biotechnology Regulatory Services
Animal and Plant Health Inspection Service
(APHIS)
United States Department of Agriculture
4700 River Road
Riverdale MD 20737, United States of America
Tel.: +1 301 734 5295
Fax: +1 301 734 3135
E-Mail: david.s.heron@aphis.usda.gov

ORGANIZATIONS

Acción Ecológica

23. Dr. Elizabeth Bravo Velasquez
 Coordinator
 Acción Ecológica
 Alejandro de Valdez
 N24-33 y La Gasca
 Quito, Ecuador
 Tel.: +593 2 547 516
 Fax: +593 2 527 583
 E-Mail: ebravo@rallt.org, ebravo@hoy.net
 Web:
www.accionecologica.org/webae/index.php

Bayer Cropscience

24. Ms. Esmeralda Prat
 Global Biosafety Manager
 Regulatory Affairs
 Bayer Cropscience
 c/o Bayer Cropscience
 Technologiemark 38
 Gent B-9052, Belgium
 Tel.: +32 9 243 0419
 Fax: +32 9 224 0694
 E-Mail: esmeralda.prat@bayercropscience.com

Federation of German Scientists

25. Dr. Ricarda Steinbrecher
 Working group member
 Working Group on Agriculture &
 Biodiversity - incl. Biotechnology and
 Biosafety
 Federation of German Scientists
 P.O. Box 1455
 Oxford Oxfordshire OX4 9BS, United
 Kingdom
 Tel.: +44 1 865 725 194
 E-Mail: r.steinbrecher@vdw-ev.de,
r.steinbrecher@gn.apc.org

Monsanto Company

26. Dr. Thomas Nickson
 Regulatory Environmental Policy
 Monsanto Company
 800 North Lindbergh Boulevard
 Saint Louis Mo 63167, United States of America
 Tel.: +314 694 2179
 Fax: +314 694 2074
 E-Mail: thomas.nickson@monsanto.com
 Web: <http://www.monsanto.com>

Public Research and Regulation Initiative

27. Dr. Piet van der Meer
 Executive Secretary
 Public Research and Regulation Initiative
 c/o Horizons sprl
 Rue d'Alaumont 16
 Lasne B-1380, Belgium
 Tel.: +32 2 652 1240
 Fax: +32 2 652 3570
 E-Mail: pietvandermeer@gmail.com

University of Canterbury

28. Prof. Jack Heinemann
 Director, Centre for Integrated Research on
 Biosafety
 School of Biological Sciences
 University of Canterbury
 Private Bag 4800
 Christchurch 8020, New Zealand
 Tel.: +643 364 2500
 Fax: +643 364 2590
 E-Mail: jack.heinemann@canterbury.ac.nz

Annex II

ACTIVITIES CARRIED OUT UNDER THE OPEN-ENDED ONLINE EXPERT GROUP ON RISK ASSESSMENT AND RISK MANAGEMENT AND BY THE AD HOC TECHNICAL EXPERT GROUP ON RISK ASSESSMENT AND RISK MANAGEMENT

Activity	Date / Location
Opening of the Online Forum and announcement of the topics and calendar of the discussion groups	6 November 2008, online
Ad hoc discussion groups under the Open-ended Online Forum on risk assessment and risk management of: (i) living modified (LM) fish; (ii) LM trees; (iii) LM microorganisms and viruses; (iv) LM pharmaplants; (v) living modified organisms (LMOs) with stacked genes or traits; (vi) post-release monitoring and long-term effects of LMOs released into the environment; and (vi) specific receiving environments; as well as on a Flowchart ("Roadmap") for risk assessment: the necessary steps to conduct risk assessment according to Annex III of the Protocol	10 November – 19 December 2008, online
First Series of Regional Real-time Online Conferences (for Europe, Latin America, Africa and Asia)	28 January – 17 February 2009, online
First Meeting of the Ad Hoc Technical Expert Group (AHTEG) on Risk Assessment and Risk Management	20 – 24 April 2009, Montreal, Canada
Meeting of the AHTEG Bureau.	24 April 2009, Montreal, Canada
Ad hoc discussion groups within the AHTEG sub-working groups for further drafting of the guidance documents	May – June 2009, online
Ad hoc discussion groups under the Open-ended Online Forum for input to the work of the AHTEG Sub-working Groups	22 June – 12 July 2009, online
Teleconference of the AHTEG Bureau	24 July 2009
Ad hoc discussion groups within the AHTEG sub-working groups for further drafting of the guidance documents and testing of the Roadmap	August – October 2009, online
Progress reports on the work of the AHTEG sub-working groups	October 2009
Meetings of the AHTEG Sub-Working Group on the Roadmap and AHTEG Bureau	12 – 14 October 2009, The Hague, Netherlands
Ad hoc discussion groups within the AHTEG sub-working groups for further drafting of the guidance documents and testing of the Roadmap	November 2009, online
Ad hoc discussion groups under the Open-ended Online Forum for further input to the work of the AHTEG sub-working groups	23 November – 14 December 2009, online

Activity	Date / Location
Ad hoc discussion group under the Open-ended Online Forum on “The way forward for the development of further guidance on risk assessment and risk management of LMOs”	7 – 14 December 2009
Ad hoc discussion groups within the AHTEG sub-working groups for further drafting of the guidance documents	January 2010, online
Second series of Regional Real-time Online Conferences (for Africa, Asia and the Pacific, WEOG and CEE, and Latin America and the Caribbean)	2-11 February 2010, online
Ad hoc discussion group under the AHTEG for final drafting of the guidance documents in preparation for the second AHTEG meeting	March 2010, online
Teleconference of the AHTEG Bureau	7 April 2010
Preparatory meetings of the AHTEG sub-working groups	19 April 2010, Ljubljana
Second meeting of the Ad Hoc Technical Expert Group	20-23 April 2010, Ljubljana

Annex III

GUIDANCE ON RISK ASSESSMENT OF LIVING MODIFIED ORGANISMS

1 This document was developed by the Ad Hoc Technical Expert Group (AHTEG) on Risk Assessment
2 and Risk Management under the Cartagena Protocol on Biosafety.²²

3 This is intended to be a “living document” that will be improved with time as new experience becomes
4 available and new developments in the field of applications of living modified organisms (LMOs) occur,
5 as and when mandated by the Parties to the Cartagena Protocol on Biosafety.

6 **PART I:**

7 **ROADMAP FOR RISK ASSESSMENT OF LIVING MODIFIED ORGANISMS**

8 This “Roadmap” provides an overview of the process of environmental risk assessment for a living
9 modified organism (LMO) in accordance with Annex III²³ to the Cartagena Protocol on Biosafety
10 (hereinafter “the Protocol”) and all other articles related to risk assessment. This Roadmap was
11 developed in response to decision BS-IV/11²⁴ of the Conference of the Parties serving as the meeting of
12 the Parties to the Protocol (COP-MOP). Annex III is the basis of the Roadmap. Accordingly, this
13 Roadmap is a guidance document and does not replace Annex III. The overall aim of the Roadmap is
14 facilitating and enhancing the effective use of Annex III by elaborating the technical and scientific
15 process of how to apply the steps and points to consider in the process of risk assessment.

16 The purpose of this Roadmap is to provide further guidance on using Annex III with additional
17 background material and links to useful references relevant to risk assessment. The Roadmap may be
18 useful as a reference for risk assessors when conducting or reviewing risk assessments and in
19 capacity-building activities.

20 The Roadmap applies to all types of LMOs²⁵ and their intended uses within the scope and objective of
21 the Protocol, and in accordance with Annex III. However, it has been developed based largely on living
22 modified crop plants because of the extensive experience to date with environmental risk assessments for
23 these organisms. It is intended to be a “living document” that will be modified and improved on over
24 time as and when mandated by COP-MOP, and in the light of new experience, information and
25 developments in the field of applications of LMOs, e.g. when other types of LMOs have been evaluated
26 more extensively in environmental risk assessments.

²² The AHTEG on Risk Assessment and Risk Management was established by the Conference of the Parties serving as the meeting of the Parties to the Cartagena Protocol on Biosafety (COP-MOP) in its decision BS-IV/11. The terms of reference for the AHTEG as set out by the Parties may be found in the annex to decision BS-IV/11 (<http://bch.cbd.int/protocol/decisions/decision.shtml?decisionID=11690>).

²³ <http://www.cbd.int/biosafety/articles.shtml?a=cpb-43> .

²⁴ <http://www.cbd.int/biosafety/cop-mop/results/?id=11690> .

²⁵ Including products thereof, as described in paragraph 5 of Annex III to the Protocol.

27 **INTRODUCTION**

28 **General introduction**

29 *Background*

30 In accordance with the precautionary approach²⁶ the objective of the Protocol is “to contribute to
31 ensuring an adequate level of protection in the field of the safe transfer, handling and use of LMOs
32 resulting from modern biotechnology that may have adverse effects on the conservation and sustainable
33 use of biological diversity, taking also into account risks to human health, specifically focusing on
34 transboundary movements”.²⁷

35 For this purpose, Parties shall ensure that risk assessments are carried out when making informed
36 decisions regarding LMOs.

37 An LMO and its use may have several effects, which may be intended or unintended, taking into account
38 that some unintended effects may be predictable. The objective of risk assessment is to *identify* and
39 *evaluate* the potential adverse effects of LMOs on the conservation and sustainable use of biological
40 diversity in the likely potential receiving environment, taking also into account risks to human health.²⁸
41 The risk assessment is performed on a case-by-case basis. What is considered an adverse effect depends
42 on protection goals and assessment end-points taken into consideration when scoping the risk assessment.
43 The choice of protection goals by the Party could be informed by Articles 7(a), 7(b) and 8(g) and
44 Annex 1 of the Convention on Biological Diversity.

45 According to the general principles of Annex III of the Protocol, risk assessments shall be based, at a
46 minimum, on information provided in accordance with Article 8 and other available scientific evidence
47 in order to identify and evaluate the possible adverse effects of LMOs on the conservation and
48 sustainable use of biological diversity, taking also into account risks to human health.²⁹

49 Annex III states that “risk assessment should be carried out in a scientifically sound and transparent
50 manner, and can take into account expert advice of, and guidelines developed by, relevant international
51 organizations. Lack of scientific knowledge or scientific consensus should not necessarily be interpreted
52 as indicating a particular level of risk, an absence of risk, or an acceptable risk. (...) Risk assessment
53 should be carried out on a case-by-case basis. The required information may vary in nature and level of
54 detail from case to case, depending on the LMO concerned, its intended use and the likely potential
55 receiving environment”.³⁰

56 *The risk assessment process*

57 Risk assessment is a structured process. Paragraph 8 of Annex III provides a description of the key steps
58 of the risk assessment process to identify and evaluate the potential adverse effects and manage risks.

²⁶ “In order to protect the environment, the precautionary approach shall be widely applied by States according to their capabilities. Where there are threats of serious or irreversible damage, lack of full scientific certainty shall not be used as a reason for postponing cost-effective measures to prevent environmental degradation” (Principle 15 of the Rio Declaration on Environment and Development) at: <http://www.unep.org/Documents/Multilingual/Default.asp?DocumentID=78&ArticleID=1163>), and in line with Articles 10.6 and 11.8 of the Protocol.

²⁷ <http://www.cbd.int/biosafety/articles.shtml?a=cpb-01> .

²⁸ Annex III, paragraph 1.

²⁹ Article 15, paragraph 1.

³⁰ Annex III, paragraphs 3, 4 and 6.

59 Paragraph 9 describes, depending on the case, points to consider in this process. The steps describe an
60 integrated process whereby the results of one step may be relevant to other steps. Also, risk assessment
61 may need to be conducted in an iterative manner, where certain steps may be repeated or re-examined to
62 increase or re-evaluate the confidence in the conclusions of the risk assessment. When new information
63 arises that could change its conclusions, the risk assessment may need to be re-examined accordingly.
64 Similarly, the issues mentioned in the ‘overarching issues’ section below can be taken into consideration
65 again at the end of the risk assessment process to determine whether the objectives and criteria that were
66 set out at the beginning of the risk assessment have been met.

67 Risk assessment is done in a comparative manner, meaning that risks associated with living modified
68 organisms should be considered in the context of the risks posed by the non-modified recipient organism
69 in the likely potential receiving environment.³¹ Additionally, experience with the same, or, as
70 appropriate, similar, genotypic or phenotypic characteristics may be taken into consideration along with
71 the non-modified recipient organism in the risk assessment of an LMO. For instance, the comparison
72 with the (near-)isogenic or closely related non-modified recipient is used in step 1 of the risk assessment
73 (see below) where the novel genotypic or phenotypic characteristics associated with the LMO are
74 identified. But when the potential consequences of adverse effects are evaluated, broader experience,
75 such as mentioned in step 3 (a), may be taken into account, when establishing a baseline. Results from
76 experimental field trials or other environmental information and experience with the same LMO may be
77 taken into account as information elements in a new risk assessment for that LMO. In all cases where
78 information, including baseline data, is derived from other sources, it is important to establish the validity
79 and relevance of the information for the risk assessment. For instance, it should be taken into account
80 that the behavior of a transgene,³² as that of any other gene, may vary because it depends on the genetic
81 and physiological background of the recipient as well as on the ecological characteristics of the
82 environment that the LMO is introduced into.

83 The concluding recommendations derived from the risk assessment in step 5 are required to be taken into
84 account in the decision-making process on an LMO. In the decision-making process, other Articles of the
85 Protocol or other relevant issues may also be taken into account and are addressed in the last paragraph
86 of this Roadmap: ‘Related Issues’.

87 A flowchart illustrating the risk assessment process according to this Roadmap is annexed hereto.

88 (*See references relevant to “[General Introduction](#)”*).

89 **Overarching issues in the risk assessment process**

90 There are some overarching issues to consider in the design/planning phase of the risk assessment
91 process to ensure the quality and relevance of the information used. These entail, among others:

- 92 • Setting criteria for relevancy in the context of a risk assessment – e.g. data may be considered
93 relevant if they can affect the outcome of the risk assessment.
- 94 • Establishment of scientifically robust criteria for the inclusion of scientific information.
 - 95 ○ Data should be of an acceptable scientific quality. Data quality should be consistent with
96 the accepted practices of scientific evidence-gathering and reporting and may include

³¹ Annex III, paragraph 5.

³² For the purpose of this document, a transgene is a nucleic acid sequence in an LMO that results from the application of modern biotechnology as described in Article 3 (i) (a) of the Protocol.

97 independent review of the methods and designs of studies. Data may be derived from a
 98 variety of sources, e.g. new experimental data as well as data from relevant peer
 99 reviewed scientific literature.

100 ○ Sound science is based on transparency, verifiability, and reproducibility (e.g. reporting
 101 of methods and data in sufficient detail, so that the resulting data and information could
 102 be confirmed independently), and on the accessibility of data (e.g. the availability of
 103 relevant, required data or information or, if requested and as appropriate, of sample
 104 material), taking into account the provisions of Article 21 of the Protocol on the
 105 confidentiality of information. The provisions of sound science serve to ensure and
 106 verify that the risk assessment is carried out in a scientifically sound and transparent
 107 manner.

108 ● Identification and consideration of uncertainty.

109 According to the Protocol, “where there is uncertainty regarding the level of risk, it may be
 110 addressed by requesting further information on the specific issues of concern or by implementing
 111 appropriate risk management strategies and/or monitoring the living modified organism in the
 112 receiving environment”.³³

113 Uncertainty is inherent in the concept of risk. To date, “there is no internationally agreed
 114 definition of ‘scientific uncertainty’, nor are there internationally agreed general rules or
 115 guidelines to determine its occurrence. Those matters are thus dealt with – sometimes differently
 116 – in each international instrument incorporating precautionary measures”.^{34, 35}

117 It should be kept in mind that uncertainty cannot always be reduced by providing additional
 118 information. For example, new uncertainties may arise as a result of the provision of additional
 119 information.

120 Considerations of uncertainty strengthen the confidence and scientific soundness of a risk
 121 assessment. In communicating the results of a risk assessment, it is important to consider and
 122 analyze in a systematic way the various forms of uncertainty that can arise at each step and in
 123 combination at step 4 of the Roadmap. An analysis of uncertainty includes considerations of its
 124 source and nature.

125 The *source(s)* of uncertainty may stem from the data/information itself and/or the choice of study
 126 design including the methods used, and the analysis of the information.

127 The *nature* of uncertainty may be described for each identified source of uncertainty arising
 128 from: (i) imperfect knowledge or lack of available information, which may be reduced with more
 129 research/information, and (ii) inherent variability.

³³ Annex III, paragraph 8 (f).

³⁴ *An Explanatory Guide to the Cartagena Protocol on Biosafety*, paragraph 57 (<http://data.iucn.org/dbtw-wpd/edocs/EPLP-046.pdf>).

³⁵ Article 10, paragraph 6, of the Protocol: “Lack of scientific certainty due to insufficient relevant scientific information and knowledge regarding the extent of the potential adverse effects of a living modified organism on the conservation and sustainable use of biological diversity in the Party of import, taking also into account risks to human health, shall not prevent that Party from taking a decision, as appropriate, with regard to the import of that living modified organism intended for direct use as food or feed, or for processing, in order to avoid or minimize such potential adverse effects.”

130 (See references relevant to "[Identification and consideration of uncertainty](#)").

131 **Context and scoping of the risk assessment**

132 In setting the context and scope for a risk assessment, a number of aspects should be taken into
133 consideration, as appropriate, that are specific to the Party involved and to the specific case of risk
134 assessment. These aspects include:

- 135 • Existing policies and strategies based on, for instance, regulations and the international
136 obligations of the Party involved; (ii) Guidelines or regulatory frameworks that the Party has
137 adopted; and (iii) Protection goals, assessment end-points, risk thresholds and management
138 strategies. Setting the context and scope for a risk assessment that are consistent with these
139 policies, strategies and protection goals may involve a process that includes risk assessors,
140 decision-makers and various stakeholders prior to conducting the actual risk assessment;
- 141 • (i) Framing the risk assessment process; (ii) Taking into account the expected (potential)
142 conditions of handling and use of the LMO; (iii) Taking into account customary practices and
143 habits that could affect the protection goals or end-points; identification of relevant questions to
144 be asked for that purpose;
- 145 • Identification of methodological and analytical requirements, including any reviewing
146 mechanisms, that is required to achieve the objective of the risk assessment as laid down, for
147 instance, in guidelines published or adopted by the Party that is responsible for conducting the
148 risk assessment (i.e. typically the Party of import according to the Protocol);
- 149 • The nature and level of detail of the information required may depend on the intended use of the
150 LMO and the likely potential receiving environment. For small scale field releases, especially at
151 early experimental stages, less information may be available compared to the information
152 available for large scale environmental release, and for commercial scale planting;
- 153 • Experience and history of use of the non-modified recipient, taking into account its ecological
154 function;³⁶ and
- 155 • Establishing criteria for describing the level of the (potential) environmental adverse effects of
156 LMOs, as well as criteria for the terms that are used to describe the levels of likelihood (step 2),
157 the magnitude of consequences (step 3) and risks (step 4) and the manageability of risks (step 5;
158 see risk assessment steps below).

159 (See references relevant to "[Context and scoping of the risk assessment](#)").

160 **THE RISK ASSESSMENT**

161 To fulfill its objective under Annex III, as well as other relevant Articles of the Protocol, risk assessment
162 is performed in five steps, as appropriate. These five steps are indicated in Paragraph 8 (a)-(e) of
163 Annex III and also detailed below. Their titles have been taken directly from the paragraphs 8 (a)-(e) of
164 Annex III.

³⁶ The term "ecological function" (or: "ecological services") provided by an organism refers to the role of the organism in ecological processes. Which ecological functions or services are taken into account here will be dependent on the protection goals set for the risk assessment. For example, organisms may be part of the decomposer network playing an important role in nutrient cycling in soils or be important as a pollen source for pollinators and pollen feeders.

165 For each step a rationale and points to consider are provided. Some points to consider are taken from
 166 paragraph 9 of Annex III, whereas others have been added based on generally accepted methodology of
 167 LMO risk assessment and risk management. The relevance of each point to consider will depend on the
 168 case being analyzed.

169 (*See references relevant to “[Risk Assessment in general](#)”*).

170 **Step 1: “An identification of any novel genotypic and phenotypic characteristics associated with**
 171 **the living modified organism that may have adverse effects on biological diversity in the likely**
 172 **potential receiving environment, taking also into account risks to human health.”³⁷**

173 *Rationale:*

174 The purpose of this step is to identify biological changes resulting from the genetic modification(s),
 175 including any deletions, compared to the non-modified organism, and identify what, if any, changes
 176 could cause adverse effects on the conservation and sustainable use of biological diversity, taking also
 177 into account risks to human health. This step is similar to the ‘hazard identification step’ in other risk
 178 assessment guidance. The comparison of the LMO is performed with the non-modified recipient, or a
 179 (near-)isogenic line or, as appropriate, with a non-modified organism of the same species, taking into
 180 consideration the new trait(s) of the LMO.

181 In this step, scientifically plausible scenarios are identified in which novel characteristics of the LMO
 182 could give rise to adverse effects in an interaction with the likely potential receiving environment. The
 183 novel characteristics of the LMO to be considered can be genotypic or phenotypic, biological. They may
 184 be intended or unintended, predicted or unpredicted. The points to consider below provide information
 185 elements on which hazard identification can be built.

186 The type and level of detail of the information required in this step may vary from case to case depending
 187 on the nature of the modification of the LMO and on the scale of the intended use of the LMO. For small
 188 scale field releases, especially at early experimental stages, less information may be available and some
 189 of the resulting uncertainty may typically be addressed by risk management measures (see step 5).

190 *Points to consider regarding the characterization of the LMO:*

- 191 (a) Relevant characteristics of the non-modified recipient (e.g. (i) its biological characteristics, in
 192 particular those that, if changed, or interacting with the new gene products or traits of the LMO,
 193 could cause changes in the behavior of the non-modified recipient in the environment in a way
 194 that may cause adverse effects; (ii) its taxonomic relationships, (iii) its origin, centers of origin
 195 and centers of genetic diversity); (iv) ecological function, and (v) as a component of biological
 196 diversity that is important for the conservation and sustainable use of the biological diversity in
 197 the context of Article 7(a) and Annex I of the Convention;
- 198 (b) Relevant characteristics of the genes and of other functional sequences, such as promoters, that
 199 have been inserted into the LMO (e.g. functions of the gene and its gene product in the donor
 200 organism with particular attention to characteristics that could cause adverse effects in the
 201 recipient);
- 202 (c) Molecular characteristics of the LMO related to the modification (e.g. (a) characteristics of the
 203 insert(s) which may include (i) gene products (intended and unintended), (ii) levels of
 204 expression, (iii) functions, (iv) insertion site in the genome of the recipient and any effects of

³⁷ The bold printed headings of each step are direct quotes from Annex III of the Protocol.

205 insertion, (v) stability or integrity within the genome of the recipient; (b) (i) the transformation
206 method, (ii) the characteristics of the vector if and, as far as it is present in the LMO, including
207 its identity, source or origin and host range) with particular attention paid to any characteristics
208 that are related to potential adverse effects. The availability and relevance of this information
209 may vary according to the type of application. Characteristics related to adverse effects may
210 also result from changed expression levels of endogenous genes due to effects of a transgene or
211 from combinatorial effects;³⁸

212 (d) Consideration of genotypic (see point to consider (c) above) and phenotypic, biological changes
213 in the LMO, either intended or unintended, in comparison with the non-modified recipient,
214 considering those changes that could cause adverse effects. These may include changes at the
215 transcriptional and translational level and may be due to the insert itself or to genomic changes
216 due to the transformation or recombination processes.

217 *Point to consider regarding the receiving environment:*

218 (e) Characteristics of the likely potential receiving environment, in particular its attributes that are
219 relevant to potential interactions of the LMO that could lead to adverse effects (see also
220 paragraph (g) below),³⁹ taking into account the characteristics that are components of biological
221 diversity;

222 (f) The intended scale and duration of the environmental release.

223 *Points to consider regarding the potential adverse effects resulting from the interaction between the*
224 *LMO and the receiving environment:*

225 (g) Characteristics of the LMO in relation to the receiving environment (e.g. information on
226 phenotypic traits that are relevant for its survival in, or its potential adverse effects on the likely
227 receiving environment – see also paragraph (e) above);

228 (h) Considerations for unmanaged and managed ecosystems (such as agricultural, forest and
229 aquaculture systems) that are relevant for the likely potential receiving environment. These
230 include the potential for dispersal of the LMO through, for instance, seed dispersal or
231 outcrossing within or between species, or through transfer into habitats where the LMO may
232 persist or proliferate;

233 (i) Potential consequences of outcrossing and flow of transgenes from an LMO to other sexually
234 compatible species, which could lead to introgression of the transgene(s) into the population of
235 sexually compatible species;

236 (j) Effects on non-target organisms;

³⁸ For the purpose of this document, the term “combinatorial effects” refers to effects that may arise from the interactions between two (or more) genes. The effects may occur at the level of gene expression, or through interactions between RNA, or among gene products. The effects may be qualitative or quantitative; quantitative effects are often referred to as resulting in antagonistic, additive or synergistic effects.

³⁹ Examples of relevant attributes of the receiving environment include, among others: (i) ecosystem type (e.g., agroecosystem, horticultural or forest ecosystems, soil or aquatic ecosystems, urban or rural environments); (ii) extension of dimension (small, medium, large or mixed scale); (iii) previous use/history (intensive or extensive use for agronomic purposes, natural ecosystem, or no prior managed use in the ecosystem); (iv) the geographical zone(s) in which the release is intended, including climatic and geographic conditions and the properties of soil, water and/or sediment; (v) specific characteristics of the prevailing faunal, floral and microbial communities including information on sexually compatible wild or cultivated species; and (vi) biodiversity status, including the status as centre of origin and diversity of the recipient organism and the occurrence of rare, endangered, protected species and/or species of cultural value.

- 237 (k) Cumulative effects;⁴⁰
- 238 (l) Effects of the incidental exposure of humans to (parts of) the LMO (e.g. exposure to pollen),
239 and the toxic or allergenic effects that may ensue;
- 240 (m) Potential adverse effects as a consequence of horizontal gene transfer (HGT) of transgenic
241 sequences from the LMO to any other organism in the likely receiving environment. With
242 regard to HGT to micro-organisms (including viruses), particular attention may be given to
243 cases where the LMO is also a micro-organism; and
- 244 (n) A consideration of uncertainty arising in step 1 that may significantly impact the identification
245 of hazards in this step (see “Identification and consideration of uncertainty” under Context and
246 scoping of the risk assessment above).

247 (*See references relevant to “[Step 1](#)”*).

248 **Step 2: “An evaluation of the likelihood of adverse effects being realized, taking into account the**
249 **level and kind of exposure of the likely potential receiving environment to the living modified**
250 **organism.”**

251 *Rationale:*

252 The potential adverse effects identified in step 1 may result in risks, but this depends on the likelihood
253 and the consequence of the effects. In order to determine and characterize the overall risk (in step 4), the
254 likelihood of each adverse effect being realized has to be assessed and evaluated beforehand.

255 One aspect to be considered is whether the receiving environment will be exposed to the LMO in such a
256 way that the identified adverse effects may actually occur, e.g. taking into consideration the intended use
257 of the LMO, and the expression level, dose and environmental fate of transgene products as well as
258 plausible pathways leading to adverse effects.

259 Other aspects to be considered here are (i) the potential of the LMO (or its derivatives resulting from
260 outcrossing) to spread and establish beyond the receiving environment (in particular into protected
261 areas), and whether that could result in adverse effects; and (ii) the possibility of occurrence of adverse
262 (e.g. toxic) effects on organisms (or on organisms other than the ‘target organism’ for some types of
263 LMOs).

264 The levels of likelihood may be expressed, for example, by the terms ‘highly likely’, ‘likely’, ‘unlikely’,
265 ‘highly unlikely’. Parties may consider describing these terms and their uses in risk assessment
266 guidelines published and/or adopted by them.

267 *Points to consider:*

- 268 (a) Information relating to the type and intended use of the LMO, including the scale and duration
269 of the release, bearing in mind, as appropriate, user habits, patterns and agronomic practices;
- 270 (b) The relevant characteristics of the likely potential receiving environment that may experience or
271 may be a factor in the occurrence of the potential adverse effects (see also step 1 (e), (f) and
272 (g)), taking into account the variability of the environmental conditions and any long-term
273 adverse effects. Levels of expression in the LMO and persistence and accumulation in the

⁴⁰ For the purpose of this document, the term “cumulative effects” refers to effects that occur due to the presence of multiple LMOs in the receiving environment.

- 274 environment (e.g. in the food chain) of substances with potentially adverse effects newly
275 produced by the LMO, such as insecticidal proteins, toxins and allergens;
- 276 (c) Available information on the location of the release and the receiving environment (such as
277 geographic and biogeographic information, including, as appropriate, coordinates, information
278 on the sexually compatible species and whether they are co-localized with the LMO and
279 whether flowering occurs at the same time, or in general, interbreeding can occur);
- 280 (d) For the case of outcrossing and outbreeding from an LMO to sexually compatible species, the
281 considerations would include: (i) the biology of the sexually compatible species; (ii) the
282 potential environment where the sexually compatible species may be located; (iii) the chance of
283 introgression of the transgene into the sexually compatible species;
- 284 (e) Expected exposure to the environment where the LMO is released and means by which
285 incidental exposure could occur at that location or elsewhere (e.g. gene flow or incidental
286 exposure due to losses during transport and handling);
- 287 (f) A consideration of uncertainty arising in step 2 (see “Identification and consideration of
288 uncertainty” under “Context and scoping of the risk assessment” above).

289 (*See references relevant to “[Step 2](#)”*).

290 **Step 3: “An evaluation of the consequences should these adverse effects be realized.”**

291 *Rationale:*

292 This step describes an evaluation of the magnitude of the consequences in the likely potential receiving
293 environment, taking into account, among others, results of tests done under different conditions such as
294 laboratory experiments or experimental field releases. The evaluation is comparative and should be
295 considered in the context of the adverse effects caused by the non-modified recipient or, if more
296 appropriate, by a near-isogenic or other non-modified organism of the same species. The evaluation may
297 also be considered in the context of the adverse effects that occur in the environment and which are
298 associated with existing practices such as various agronomic practices, for example, for pest or weed
299 management if such information is available and relevant. The evaluation of the consequence of adverse
300 effects may be expressed as, for instance, ‘major’, ‘intermediate’, ‘minor’ or ‘marginal’. Parties may
301 consider describing these terms and their uses in risk assessment guidelines published and/or adopted by
302 them.

303 *Points to consider:*

- 304 (a) Relevant experience with the consequences of existing practices with the non-modified
305 recipient or, if more appropriate, with a non-modified organism of the same species in the likely
306 potential receiving environment, may be useful in order to establish baselines to evaluate, for
307 example, the consequences of (i) agricultural practices, such as the level of inter- and intra-
308 species gene flow, dissemination of the recipient, abundance of volunteer plants in crop
309 rotation; occurrence of pests and/or beneficial organisms such as pollinators and pest predators;
310 or (ii) pest management, including effects on non-target organisms in pesticide applications
311 while following accepted agronomic practices;
- 312 (b) Adverse effects which may be direct and indirect, immediate and delayed. Some of these
313 adverse effects may result from combinatorial and cumulative effects;
- 314 (c) Results from laboratory experiments examining, *inter alia*, dose-response relationships (e.g.,
315 EC 50s, LD 50s) and from field trials evaluating, for instance, potential invasiveness;

316 (d) For the case of outcrossing to sexually compatible species, the possible adverse effects that may
317 occur, after introgression, due to the expression of the transgenes in the sexually compatible
318 species; and

319 (e) A consideration of uncertainty arising in step 3 that may significantly impact the evaluation of
320 consequences should the adverse effects be realized (see “Identification and consideration of
321 uncertainty” under Context and scoping of the risk assessment above).

322 (*See references relevant to “[Step 3](#)”*).

323 **Step 4: “An estimation of the overall risk posed by the living modified organism based on the**
324 **evaluation of the likelihood and consequences of the identified adverse effects being realized.”**

325 *Rationale:*

326 The purpose of this step is to determine and characterize the level of the overall risk based on the
327 identified individual risks posed by the LMO on the conservation and sustainable use of biological
328 diversity, taking also into account human health. The individual risks are determined on the basis of an
329 analysis of the potential adverse effects identified in step 1, their likelihood (step 2) and consequences
330 (step 3), and also taking into consideration any relevant uncertainty that emerged in the preceding steps.

331 It should then be determined whether the assessed risks meet the criteria set out in the protection goals,
332 assessment endpoints and thresholds, as established in relevant legislation of the Party or in its practice.
333 Where there is uncertainty regarding the level of risk, it may be addressed by requesting further
334 information on the specific issues of concern or by implementing appropriate risk management strategies
335 and/or monitoring the LMO in the receiving environment (see also step 5). Description of the risk
336 characterization may be expressed as, for instance, ‘high’, ‘medium’, ‘low’, ‘negligible’ or
337 ‘indeterminate due to uncertainty or lack of knowledge’. Parties may consider describing these terms and
338 their uses in risk assessment guidelines published and/or adopted by them.

339 To date, there is no universally accepted method to estimate the overall risk but rather a number of
340 methods are available for this purpose. The outcome of this step may be, for example, a description
341 explaining how the estimation of the overall risk was performed.

342 *Points to consider:*

343 (a) The identified potential adverse effects (step 1);

344 (b) The assessments of likelihood (step 2);

345 (c) The evaluation of the consequences (step 3);

346 (d) Any interaction between the identified individual risks;

347 (e) Any cumulative effect due to the presence of multiple LMOs in the receiving environment; and

348 (f) A consideration of uncertainty arising in this and the previous steps (see “Identification and
349 consideration of uncertainty” under Context and scoping of the risk assessment above).

350 (*See references relevant to “[Step 4](#)”*).

351 **Step 5: “A recommendation as to whether or not the risks are acceptable or manageable,**
352 **including, where necessary, identification of strategies to manage these risks”**

353 *Rationale:*

354 In this way, step 5 provides an interface between the process of risk assessment and the process of
355 determining whether risk management measures are necessary and, if so, which measures could be
356 implemented to manage the risks associated with the LMO.

357 The evaluation of the overall risk on the basis of the identified individual risks conducted in the previous
358 step may lead to the conclusion that the identified risks are not acceptable in relation to the established
359 protection goals, assessment end-points and risk thresholds, also when taking into account risks posed by
360 the non-modified recipient and its use. Then the question arises whether risk management options can be
361 identified that have the potential to remove the identified risks or reduce their magnitude. In the process
362 of the formulation of risk management options, the effect of the proposed options on the identified risks
363 should be explained. The appropriate steps of the risk assessment should then be reiterated by taking into
364 account the implementation of the risk management options to estimate the new levels of likelihood,
365 consequence and risk and to assess if the risk management measures are appropriate and sufficient.

366 The issues mentioned in the ‘overarching issues’ section can be taken into consideration again at the end
367 of the risk assessment process to evaluate whether the objectives and criteria that were set out at the
368 beginning of the risk assessment have been met.

369 The recommendation of acceptability of risk(s) should acknowledge the previously identified
370 uncertainties. Some uncertainties may be reduced by monitoring (e.g. checking the validity of
371 assumptions about the ecological effects of the LMO), requests for more information, or implementing
372 the appropriate risk management options.

373 The recommendation(s) as to whether or not the risks are acceptable or manageable and
374 recommendations for risk management options are submitted for consideration in the decision-making
375 process.

376 *Points to consider related to the acceptability of risks:*

377 (a) The criteria for the establishment of acceptable/unacceptable levels of risk, including those set
378 out in national legislation or guidelines, as well as the protection goals of the Party, as
379 identified when setting the context and scope for a risk assessment;

380 (b) In establishing a baseline for the comparison of the LMO, any relevant experience with the use
381 of the non-modified recipient, and practices associated with its use in the potential receiving
382 environment; and

383 (c) The feasibility of the adoption of risk management or monitoring strategies.

384 *Points to consider related to the risk management strategies:*

385 (d) Existing management practices, if applicable, that are in use for the non-modified recipient
386 organism or for other organisms that require comparable risk management and that might be
387 appropriate for the LMO being assessed, e.g. isolation distances to reduce outcrossing potential
388 of the LMO, modifications in herbicide or pesticide management, crop rotation, soil tillage,
389 etc.;

390 (e) Methods to detect and identify the LMO and their specificity, sensitivity and reliability in the
391 context of environmental monitoring (e.g. monitoring for short- and long-term, immediate and
392 delayed effects; specific monitoring on the basis of scientific hypotheses and supposed
393 cause/effect relationship as well as general monitoring) including plans for appropriate
394 contingency measures to be applied in case the results from monitoring call for them;

395 (f) Management options in the context of the intended use (e.g. mitigating the effect of an LMO
396 producing insecticidal proteins by the use of refuge areas to minimize the development of
397 resistance against these proteins).

398 (*See references relevant to “[Step 5](#)”*).

399 **RELATED ISSUES**

400 Some members of the AHTEG considered some issues to be related to risk assessment and decision-
401 making process but outside the scope of this Roadmap. These issues were, *inter alia*:

- 402 • Risk management (Article 16);
- 403 • Capacity-building (Article 22);
- 404 • Public awareness and participation (Article 23);
- 405 • Socio-economic considerations (Article 26);
- 406 • Liability and redress (Article 27);
- 407 • Co-existence;
- 408 • Ethical issues.

Annex

FLOWCHART FOR RISK ASSESSMENT

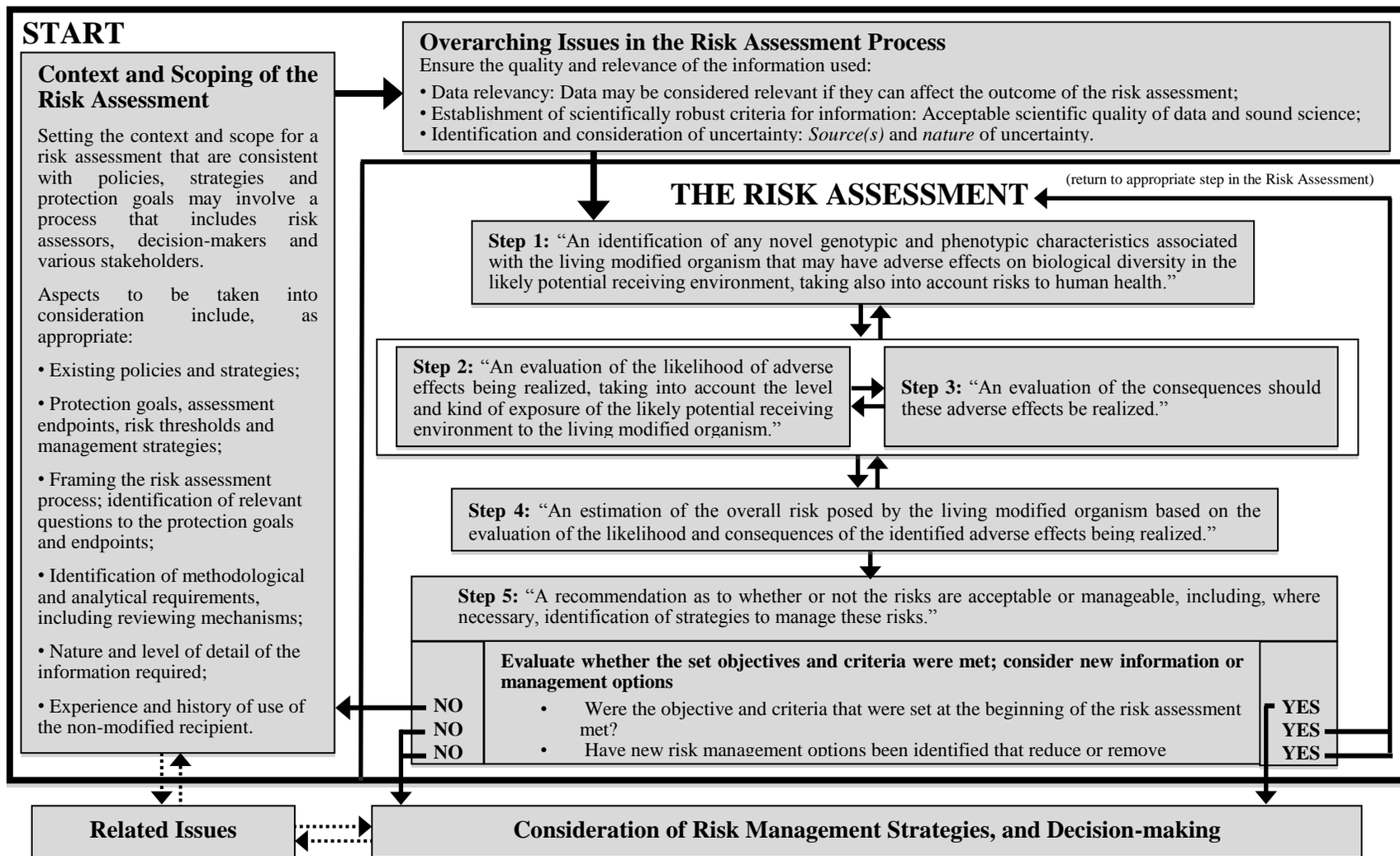


Figure 1. The Roadmap for Risk Assessment. The flowchart represents the steps to *identify* and *evaluate* the potential adverse effects of LMOs on the conservation and sustainable use of biological diversity in the likely potential receiving environment, taking also into account risks to human health. The box around steps 2 and 3 shows that these steps may sometimes be considered simultaneously or in reverse order.

PART II

SPECIFIC TYPES OF LMOs AND TRAITS

A. RISK ASSESSMENT OF LIVING MODIFIED ORGANISMS WITH STACKED GENES OR TRAITS

INTRODUCTION

Worldwide, a growing number of LMOs with stacked transgenic traits, particularly LM crops, are being developed for commercial uses. As a result, the number of stacked genes in a single LMO and the number of LMOs with two or more transgenic traits is growing.

Stacked transgenic traits can be produced through different approaches. In addition to the cross-hybridising of two LMOs, multiple trait characters can be achieved by transformation with a multigene cassette, retransformation of an LMO or simultaneous transformation with different transgene cassettes (i.e., cotransformation).

This guidance document focuses on stacked transgenic traits that have been produced through cross-breeding of two or more LMOs.

LMOs with multiple transgenic traits resulting from re-transformation, co-transformation or transformation with a multigene cassette should be assessed according to the Roadmap.

This guidance document complements the Roadmap for Risk Assessment developed by the AHTEG on Risk Assessment and Risk Management, and focuses on issues that are of particular relevance to the risk assessment of LMOs with stacked events generated through cross breeding of single or multiple event LMO.

This is intended to be a “living document” that will be shaped and improved with time as new information and/or experience becomes available and new developments in the field of applications of LMOs occur, as and when mandated by the Parties to the Protocol.

OBJECTIVE

The objective of this document is to give additional guidance on the risk assessment (RA) of LMOs with stacked events generated through conventional crossing of single or multiple event LMOs. Accordingly, it is meant to complement the Roadmap for Risk Assessment⁴¹ and address special aspects of LMOs with stacked transgenes/traits resulting from the conventional crossing. For the time being it will be restricted to plant LMOs.⁴²

⁴¹ In accordance with a mandate from the Parties to the Cartagena Protocol on Biosafety (the Protocol), the AHTEG has developed ‘a “roadmap”, such as a flowchart, on the necessary steps to conduct a risk assessment in accordance with Annex III to the Protocol and, for each of these steps,’ has provided ‘examples of relevant guidance documents’. The Roadmap is presented, together with the present document, to the Parties of the Protocol on the occasion of the fifth meeting of the Conference of the Parties serving as the meeting of the Parties.

⁴² It is also restricted to those LMO generated through the methods of Modern Biotechnology as defined in Art. 3 (i) (a) of the Protocol. LMOs derived from fusion of cells are not covered in this document.

USE OF TERMS

Transformation event (TraEv)

For the purpose of this document, a transformation event (TraEv) is an LM plant which results from the use of modern biotechnology applying *in vitro* nucleic acid techniques⁴³ that may involve, but is not limited to, single or multiple gene transformation cassettes. In either case, the result will be one transformation event.

Stacked event (StaEv)

For the purpose of this document, a stacked event (StaEv) is an LM plant generated through conventional cross breeding of two or more single parental transformation events (TraEvs) or two already stacked events. Accordingly the transgene⁴⁴ cassettes may be physically unlinked (i.e. located separately in the genome) and may segregate independently.

Unintentional stacked event

Unintentional stacked events are the result of outcrossing of stacked events into other LMOs or compatible relatives in the receiving environment. Depending on the segregation pattern of the stacked genes this may result in new and/or different combinations of TraEvs.

SCOPE

This guidance document focuses on stacked events (StaEv) resulting from conventional crossings between two or more single transformation events (TraEv) as parental lines so that the resulting LMO contains two or more transgenic traits. It is understood that the individual TraEvs making up the StaEv have been assessed previously in accordance with Annex III of the Cartagena Protocol on Biosafety and as described in the Roadmap.

ISSUES TO BE CONSIDERED IN THE RISK ASSESSMENT

Assessment of sequence characteristics at the insertion sites and genotypic stability (*see step 1, Point to consider (c) of the Roadmap for Risk Assessment*)

Rationale:

Although recombination, mutation and rearrangements are not limited to LMOs, the combination of transgenic traits via cross breeding may further change the molecular characteristics of the inserted genes/gene fragments at the insertion site and/or influence the regulation of the expression of the transgenes. In addition, changes to the molecular characteristics may influence the ability to detect the LMO, which may be needed in the context of risk management measures (see step 5 of the Roadmap). The reappraisal of the molecular sequence at the insertion sites, and the intactness of the transgenes may be confirmative to the molecular characteristics of the parental LMOs, but may also be a basis for assessing any intended or unintended possibly adverse effects on the conservation and sustainable use of biological diversity in the likely potential receiving environment and of potential adverse effects on

⁴³ See Article 3 (i) (a) of the Protocol.

⁴⁴ For the purpose of this document, a transgene is a nucleic acid sequence that results from the application of modern biotechnology as described in Article 3 (i) (a) of the Protocol.

human health. The extent of the reexamination may vary case by case and take into account the results of the parental LMO risk assessment.

Assessment of potential interactions between combined events and the resulting phenotypic effects
(see step 1, point to consider (d) of the Roadmap for Risk Assessment)

Rationale:

The combination of two or more TraEvs resulting in a StaEv may influence the expression level of each of the transgenes and there may be interaction between the genes and the expressed products of the different transgenes. In addition, the stacked transgenes may alter the expression of endogenous genes.

Therefore, in addition to information about the characteristics of the parental single-TraEv LMOs, specific information on potential for interactions between the altered or inserted genes, stacked proteins or modified traits and endogenous genes and their products in the StaEv LMO should be considered and assessed. For example, it should be assessed whether the different transgenes affect the same biochemical pathways or physiological processes, or are expected to or may have any combinatorial effects that may result in potential for new or increased adverse effects relative to the parent LMOs.

Assessment of combinatorial and cumulative effects of stacked event LMOs on the conservation and sustainable use of biological diversity in the likely potential receiving environment, taking also into account potential adverse effects to human health (see step 1, point to consider (c), step 2, point to consider (c) and step 3, point to consider (b) of the Roadmap for Risk Assessment)

Rationale:

Assessment of combinatorial and cumulative effects⁴⁵ is based on the environmental risk assessment data for the StaEv LMO in comparison to the closely related non-modified recipient species and the parent LMOs in the likely receiving environment, taking into consideration the results of the genotypic and phenotypic assessments outlined above.

If potential new or increased adverse effects on the conservation and sustainable use of biological diversity or on human health are identified in relation to the StaEv through the above analysis of possible interactions, additional supporting data on StaEv may be required, such as:

- (a) Phenotypic characteristics, including the levels of expression of any introduced gene products or modified traits, compared to the parent LMOs and to relevant non-modified recipient organisms (plants);
- (b) Compositional analysis (e.g. levels of expression in the LMO and persistence and accumulation in the environment, such as in the food chain) of substances with potentially harmful effects newly produced by the StaEv, (e.g. insecticidal proteins, allergens, anti-nutritional factors, etc.) in amounts that differ from those produced by the parental LMOs or non-modified recipient organisms;
- (c) Additional information depending on the nature of the combined traits. For example, further toxicological analysis of the StaEv may be required to address any combinatorial effects arising from the stacking of two or more insecticidal traits that result in a broadened target range or increased toxicity.

⁴⁵ See definition of combinatorial and cumulative effects in the Roadmap (footnotes 38 and 40, respectively).

Also, indirect effects due to changed agricultural management procedures, combined with the use of the transgenic stacked event LMO, should be taken into consideration.

Intentional and unintentional StaEvs may have altered environmental impacts as a result of cumulative and combinatorial effects of the stacked traits prevalent in different LMOs of the same species in the receiving environment. Unintentional StaEvs may arise from outcrossing with other LMOs of the same species or cross-compatible relatives (see “Use of terms”). If a number of different StaEvs are cultivated in the same environment a number of varying unintentional StaEvs may occur. Changed impacts on non-target organisms or a change in the range of non-target organisms in the likely receiving environment should be taken into account.

Development of specific methods for distinguishing the combined transgenes in a stacked event from the parental LMOs (*see step 5, point to consider (d) of the Roadmap for Risk Assessment*)

Rationale:

Some of the risk management strategies for StaEvs may involve methods for the detection and identification of these LMOs in the context of environmental monitoring. Currently, many detection methods for LMOs rely on DNA-based techniques, such as polymerase chain reaction (PCR) or protein based ELISA tests targeted to single transformation events. The methods used to detect the transgene in the parental lines may not be sensitive or specific enough to differentiate between single parental transformation events and the same event being part of a stacked event. A special problem may arise particularly in the cases where the StaEv contains multiple transgenes with similar DNA sequences. Therefore, the detection of each and all individual transgenes in a StaEv may become a challenge and need special consideration.

BIBLIOGRAPHIC REFERENCES

See references relevant to the “[*Guidance Document on Risk Assessment of LMOs with Stacked Genes or Traits*](#)”.

B. RISK ASSESSMENT OF LIVING MODIFIED CROPS WITH TOLERANCE TO ABIOTIC STRESS

1 INTRODUCTION

2 The aim of this document is to provide further guidance for the risk assessment of living modified (LM)
3 crops with improved tolerance to abiotic stress.

4 This guidance document should be considered in the context of the Cartagena Protocol on Biosafety. The
5 elements of Articles 15 Annex III of the Protocol also apply to LM crops with tolerance to abiotic stress.
6 Accordingly, the methodology and points to consider⁴⁶ contained in Annex III are also applicable to this
7 type of LMO.

8 The potential environmental adverse effects of an LM crop with abiotic stress tolerance depends on (i)
9 the receiving environment; (ii) the modified crop, (iii) phenotypic changes resulting from the genotypic
10 changes made to the plant and (iv) its intended use. A risk assessment would be performed on a case-by-
11 case basis in accordance with Annex III of the Protocol.

12 This guidance document complements the Roadmap for Risk Assessment developed by the AHTEG on
13 Risk Assessment and Risk Management, and focuses on issues that are of particular relevance to the risk
14 assessment of LM crops tolerant to abiotic stress.

15 USE OF TERMS

16 “Abiotic stresses” are environmental conditions caused by non-living factors that are detrimental or
17 suboptimal to the growth, development and/or reproduction of a living organism. Types of abiotic
18 stresses include, for example, drought, salinity, cold, heat, soil pollution and air pollution (e.g., nitrous
19 oxides, ozone).

20 RISK ASSESSMENT

21 While the same general principles used in the risk assessments of other types of LMOs also apply to LM
22 crops with increased tolerance to abiotic stress, there are a number of specific issues that may be of
23 particular importance when assessing the risks of LM crops tolerant to abiotic stresses.

24 Questions that may be relevant to the risk assessment of LM crops with tolerance to abiotic stress in
25 connection with the intended use and receiving environment include:

- 26 • Would the tolerance trait have the potential to increase the invasiveness, persistence or
27 weediness of the LM crop that causes adverse effects to other organisms?
- 28 • Would a LM plant expressing tolerance to a particular abiotic stress have other advantages in
29 the targeted receiving environment that cause adverse effects?
- 30 • Would any LMO arising from outcrossing with the abiotic stress tolerant LM crop, have the
31 potential to colonize an ecosystem beyond the targeted receiving environment?
- 32 • Would the abiotic stress tolerance trait, for example, via pleiotropic effects, have the potential to
33 affect, *inter alia*, pest and disease resistance mechanisms of the LM crop?

⁴⁶ Paragraphs 8 and 9 of Annex III, respectively.

34 Some of the potential adverse effects to be evaluated in the risk assessment, from the introduction of
35 crops tolerant to abiotic stress into the environment include, for example: a) increased selective
36 advantage(s) other than the intended tolerance trait; b) increased persistence in agricultural areas and
37 increased invasiveness in natural habitats; c) adverse effects on organisms exposed to the crop; and d)
38 consequences of potential gene flow to wild or conventional relatives. While these adverse effects may
39 exist regardless of whether the tolerant crop is a product of modern biotechnology or conventional
40 breeding, some specific issues may be more relevant in the case of abiotic stress tolerant LM crops.

41 **Characterization of the LM crop with tolerance to abiotic stress in comparison with its non-**
42 **modified crop** (*see step 1 of the Roadmap for Risk Assessment*)

43 *Rationale:*

44 The first step in the risk assessment process involves the characterization of genotypic or phenotypic,
45 biological, intended and unintended changes associated with the abiotic stress tolerant LM crop that may
46 have adverse effects on biodiversity in the likely receiving environment, taking into account risks to
47 human health. This step is the ‘hazard identification step’ in other risk assessment guidance.

48 The identification of genotypic and phenotypic changes in the abiotic stress tolerant LM crop, either
49 intended or unintended, is typically done in comparison with the non-modified recipient organism (see
50 step 1 of the Roadmap). The non-modified comparator provides the baseline information for comparison
51 of trials when it is grown at the same time and location as the LM crop. Comparisons with the observed
52 range of changes in the non-modified crop in different environments, also provides baseline information.

53 *Challenges with respect to experimental design:* Abiotic stress crops may present unique challenges in
54 experimental design for risk assessment. In some cases, for instance, an approach uses different
55 reference plant lines, which typically include a range of genotypes representative of the natural variation
56 in the crop species. In such conditions, choosing appropriate comparators could be a challenge and there
57 are several proposals on whether and how the comparative approach can be used to characterize LM
58 crops tolerant to abiotic stress in these likely receiving environments. Another important consideration is
59 whether the experimental design properly controlled for the effect of the abiotic stress trait. In the
60 extreme case, when the non-modified crop has never been grown in the range of conditions of the
61 receiving environment because the abiotic stress conditions prevent or severely affect the growth of the
62 non-modified crop, a comparative approach between the LM crop and the non-modified crop will need to
63 be adjusted.

64 The use of non-isogenic reference lines can make it more difficult to identify statistically meaningful
65 differences. In some situations when a comparator may not be available to carry out a meaningful
66 comparison, a characterization of the abiotic stress tolerant LM crop as a novel genotype in the receiving
67 environment may be conducted. In the future, information available from “omics” technologies, for
68 example, “transcriptomics” and “metabolomics”, if available, may help to detect phenotypes (e.g., the
69 production of a novel allergen or anti-nutrient) that cannot be detected using a comparison between field
70 grown plants at a suboptimal condition.

71

72 *Points to consider:*

- 73 (a) Characteristics of the LM crop under the abiotic stress and non-stress conditions and under
74 different stresses, if applicable;
- 75 (b) Likelihood of gene flow to wild or domestic relatives; and
- 76 (c) Whether one or more suitable comparators are available and the possibility of their use in the
77 appropriate experimental design.

78 **Unintended characteristics** (*see step 1 of the Roadmap for Risk Assessment*)79 *Rationale:*

80 Both intended and unintended changes to the LM crop which are directly or indirectly associated with the
81 abiotic stress tolerance that may have adverse effects should be identified. These include changes to the
82 biology of the crop plant (e.g. if the genes alter multiple characteristics of the plant) or to its distribution
83 range in relation to the potential receiving environment (e.g. if the plant can grow where it has not grown
84 before), that may cause adverse effects.

85 The abiotic-stress-tolerant LM crop may have unintended characteristics such as tolerances to other types
86 of biotic and abiotic stresses, which could lead to a selective advantage of these crop plants under
87 conditions other than that related to the modified trait. For instance, crops modified to become tolerant to
88 drought or salinity may be able to compete better than their counterparts at lower and higher growing
89 temperatures.

90 It is also possible the LM crops with enhanced tolerance to an abiotic stress could have changes in seed
91 dormancy, viability, and/or germination rates under other types of stresses. Particularly if genes involved
92 in abiotic stress are also involved in crucial steps in physiology, modifications involving these genes
93 may, therefore, have pleiotropic effects. Such LM crops may also transfer genes for stress tolerance at
94 higher frequencies than observed in non-modified crops.

95 A potential mechanism for interactions between abiotic and biotic stresses may exist in plants. For
96 example, drought or salinity-tolerant LM crops may acquire a changed tolerance to biotic stresses, which
97 could result in changed interactions with their predators, parasitoids and pathogens, and, therefore, have
98 both direct and indirect effects on organisms that interact with them.

99 *Points to consider:*

- 100 (a) Any intended or unintended change that may lead to selective advantage or disadvantage
101 acquired by the LM crop under other abiotic or biotic stress conditions that could cause
102 adverse effects;
- 103 (b) Any change in the resistance to biotic stresses and how these could affect the population of
104 organisms interacting with the LM crop; and
- 105 (c) A change in the substances (e.g., toxin, allergen, or nutrient profile) of the LM crop that could
106 cause adverse effects.

107 **Increased persistency in agricultural areas and invasiveness of natural habitats** (*see steps 1, 3 and 5
108 of the Roadmap for Risk Assessment*)

109 *Rationale:*

110 Climate change, water depletion or elevated salt content are examples of factors that limit the growth,
111 productivity, spread or persistence of a crop. Expression of the genes for abiotic stress tolerance could
112 result in increased persistence of the modified crop in agricultural areas. Expression of these genes may
113 also alter the capacity of LM crops to spread to and establish in climatic and geographic zones beyond
114 those initially considered as the likely or potential receiving environments.

115 The gene(s) inserted for tolerance to, for instance, drought and salinity might also affect molecular
116 response mechanisms to other forms of abiotic stress, such as cold temperatures. For example, when the
117 genetic modification affects genes that also regulate key processes in seeds, such as abscisic acid (ABA)
118 metabolism, physiological characteristics such as dormancy and accumulation of storage lipids may also
119 be changed. In such cases, the seeds of a tolerant crop, modified for drought or salinity tolerance, may
120 acquire in addition tolerance to cold resulting in an increased winter survivability of the seeds. Therefore,
121 an abiotic stress-tolerant crop may acquire the potential to persist better than its conventional counterpart
122 under different abiotic-stress conditions.

123 *Points to consider:*

- 124 (a) Consequences of the increased potential for persistency of the modified crop in agricultural
125 habitats and consequences of increased potential for invasiveness in natural habitats;
- 126 (b) Need for control measures if the abiotic stress-tolerant crop shows a higher potential for
127 persistency in agricultural or natural habitats, that could cause adverse effects;
- 128 (c) Characteristics that are generally associated with weediness such as prolonged seed dormancy,
129 long persistence of seeds in the soil, germination under a broad range of environmental
130 conditions, rapid vegetative growth, short lifecycle, very high seed output, high seed dispersal
131 and long-distance seed dispersal; and
- 132 (d) Effects of climate change on agriculture and biodiversity and how this could change the habitat
133 range of the LM crop in comparison to the non modified crop.
- 134 (e) If the LM crop expressing tolerance, would have a change in its agriculture practices.

135 **BIBLIOGRAPHIC REFERENCES**

136 *See references relevant to the “[Guidance Document on Risk Assessment of LM Crops with Tolerance to](#)*
137 *[Abiotic Stress](#)”.*

C. RISK ASSESSMENT OF LIVING MODIFIED MOSQUITOES

1 INTRODUCTION

2 Living modified (LM) mosquitoes are being developed through modern biotechnology to reduce
3 transmission of vector borne human pathogens, particularly those that cause malaria, dengue and
4 chikungunya. Control, including eradication of such diseases, is a recognized public health goal. Some of
5 the strategies being developed are to control mosquito vectors by suppressing their population or
6 reducing their competence. These strategies can be subcategorized according to the technology involved
7 and the method used. Some are intended to develop LM mosquitoes that are genetically modified to be
8 sterile or self-limiting (i.e., unable to pass the modified trait on indefinitely through subsequent
9 generations). Modern biotechnology techniques for developing sterile LM mosquitoes are different from
10 those based on the use of irradiation to induce male sterility.

11 Other modern biotechnology strategies are also being used for developing LM mosquito populations that
12 are self-sustaining or self-propagating (i.e., heritable modifications intended to spread through the target
13 population). The strategy used is an important factor to be considered in the risk assessment and risk
14 management process since there might be different points to be considered, depending on the specific
15 strategy used.

16 The biology and ecology of mosquitoes on the one hand, and their impact on public health as vectors of
17 human and animal diseases on the other hand, pose new considerations and challenges during the risk
18 assessment process, which have mainly dealt with LM crop plants thus far.

19 This guidance document provides information for the risk assessment of environmental releases of LM
20 mosquitoes and aims at helping to conduct risk assessments for environmental releases of LM
21 mosquitoes. Although the focus of this guidance is on LM mosquitoes, in principle, it may also be useful
22 for the risk assessment of similar non-LM mosquito strategies.

23 The main emphasis of this guidance document is the assessment of potential risks to biodiversity.
24 Nevertheless, the potential adverse effects to human health arising from environmental releases of LM
25 mosquitoes should also be considered.

26 This guidance document complements the Roadmap for Risk Assessment developed by the AHTEG on
27 Risk Assessment and Risk Management and focuses on specific issues that may need special
28 consideration on the risk assessment for environmental releases of LM mosquitoes.

29 OBJECTIVE

30 The objective of this document is to give additional guidance on the risk assessment (RA) of LM
31 mosquitoes in accordance with Annex III to the Cartagena Protocol on Biosafety.⁴⁷ Accordingly, it aims
32 at complementing the Roadmap for Risk Assessment on specific issues that may need special
33 consideration for the environmental release of LM mosquitoes.

⁴⁷ The Parties to the Cartagena Protocol on Biosafety have mandated the AHTEG to ‘develop a “roadmap”, such as a flowchart, on the necessary steps to conduct a risk assessment in accordance with Annex III to the Protocol and, for each of these steps, provide examples of relevant guidance documents’. The Roadmap is meant to provide reasoned guidance on how, in practice, to apply the necessary steps for environmental risk assessment as set out in Annex III of the Protocol. The Roadmap also demonstrates how these steps are interlinked.

34 **SCOPE**

35 This document focuses on the specific aspects of risk assessment of LM mosquitoes developed to be
36 used in the control of human and zoonotic diseases such as malaria, dengue, chikungunya, yellow fever
37 and West Nile.

38 **ISSUES TO BE CONSIDERED IN THE RISK ASSESSMENT**

39 *(See step 1 of the Roadmap for Risk Assessment of LMOs)*

40 Specific and comprehensive considerations should be undertaken with respect to the potential adverse
41 effects of a particular LM mosquito, taking into account the species of the mosquito, the LM trait, the
42 intended receiving environment, and the objective and scale of the intended release. These considerations
43 should focus on, for instance: (a) description of the genetic modification; (b) the kinds of possible
44 adverse effects for which there are scientifically plausible scenarios; (c) the species and ecological
45 processes that could be affected by the introduction of the LM mosquitoes; (d) the protection goals of the
46 country where the LM mosquitoes will be introduced; and (e) a conceptual link between the identified
47 protection goals and the introduction of the LM mosquito into the environment.

48 The biology and, to some extent, the ecology of the mosquito species that transmit malaria and dengue
49 are well known in many regions of the world. However, in certain regions and in the environment where
50 the LM mosquito is likely to be released, more information may be needed depending on the nature and
51 scale of the LM strategy to be deployed. In many of these environments few studies have been conducted
52 to examine gene flow among vectors, their mating behaviour, the interactions between vectors sharing
53 one habitat, how pathogens respond to the introduction of new vectors, etc. Such information may be
54 needed to establish a baseline in order to successfully assess the risks of LM mosquitoes. Additionally,
55 methods for the identification of specific ecological or environmental hazards are also needed.

56 **Effects on biological diversity (species, habitats, ecosystems, and ecosystem services)**

57 *(See step 2 of the Roadmap for Risk Assessment of LMOs)*

58 *Rationale:*

59 The release of LM mosquitoes may have a negative impact on the target vector and pathogen⁴⁸ and other
60 species, such as:

61 *New or more vigorous pests, especially those that have adverse effects on human health:* (i) the released
62 LM mosquitoes may not function as expected, for example gene silencing or production failures could
63 result in the release of non-sterile or competent mosquitoes and thus increase the vector population or
64 disease transmission; (ii) the released LM mosquitoes could transmit another disease more efficiently
65 than indigenous non-LM mosquitoes, such diseases might include yellow fever, chikungunya, etc.; (iii)
66 suppression of the target mosquito might result in the population of another vector species to increase
67 and result in higher levels of the target disease or the development of a new disease in humans and/or
68 animals. These other vector species may include other mosquito vectors of other diseases; (iv) the
69 released LM mosquitoes might become pests; (v) the released LM mosquitoes might cause other pests to
70 become more serious, including agricultural pests and other pests that affect human activities.

⁴⁸ For the purpose of this guidance, the term “target vector” refers to the mosquito that transmits the disease and “target pathogen” is the disease causing agent transmitted by the target mosquito.

71 *Harm to or loss of other species:* The released LM mosquitoes might cause other species (for instance
72 fish that rely seasonally on mosquitoes for food) to become less abundant. These include species of
73 ecological, economic, cultural and/or social importance such as wild food, endangered, keystone, iconic
74 and other relevant wildlife species. Ecological effects might result from competitive release if the target
75 mosquito population is reduced or from trophic consequences of species that rely on mosquitoes for food
76 at specific times of the year. Effects may also occur if (i) the target mosquitoes transmit a disease to
77 animal species, (ii) the released LM mosquitoes transmit a disease to animal species more efficiently,
78 (iii) another vector of an animal disease was released from control when the target mosquito population
79 was reduced, or (iv) the population of a target pathogen is reduced or lost and this may affect other
80 organisms that interact with it.

81 Although mosquitoes, like other insects, typically have strong reproductive isolating mechanisms that
82 will not allow interspecific gene flow, if sterile interspecific mating between released LM mosquitoes
83 and other mosquito species should occur, it could disrupt the population dynamics of these other species,
84 leading to harm or loss of valued ecological species. Moreover, cessation of transmission of pathogens to
85 other animals (e.g., West Nile virus to birds, Rift Valley fever virus to African mammals) might alter the
86 population dynamics of those species, favouring increases in their numbers.

87 *Disruption of ecological communities and ecosystem processes:* The ecological communities in the
88 ephemeral, small aquatic habitats occupied by the non-LM mosquitoes are unlikely to be disrupted
89 beyond the possibilities already addressed above under “harm to or loss of other species.” However, if
90 the released LM mosquitoes were to inhabit natural habitats (e.g. tree-holes), disruption of the associated
91 community is a possibility. The released LM mosquitoes might degrade some valued ecosystem process.
92 This might include processes such as pollination or support of normal ecosystem functioning. These
93 processes are often referred to as “ecosystem services”. However, the valued ecosystem processes may
94 also be culturally or socially specific. Under some circumstances, mosquito species are significant
95 pollinators. In those cases, mosquito control of any kind might reduce the rate of pollination of some
96 plant species or cause a shift to different kinds of pollinators. Habitats in which mosquitoes are the
97 dominant insect fauna (e.g., high Arctic tundra, tree holes) would be changed if mosquitoes were
98 eliminated; however, the common target vector species are usually associated with human activity and
99 therefore not as closely tied to ecosystem services.

100 *Points to consider:*

- 101 (a) Impacts on the target mosquitoes and pathogens resulting from the use of the strategy under
102 consideration;
- 103 (b) Whether the LM mosquitoes have the potential of causing adverse effects on other species
104 which will result in the other species becoming agricultural, aquacultural, public health or
105 environmental pests, or nuisance or health hazards;
- 106 (c) Whether the target mosquito species is native or invasive to a given area;
- 107 (d) The habitat range of the target mosquito species and whether the habitat range is likely to be
108 affected by climate change;
- 109 (e) Any other species (e.g. animal hosts, larval pathogens or predators of mosquitoes) in addition to
110 the pathogen, that typically interact with the LM mosquito in the likely receiving environment;
- 111 (f) Whether the release of LM mosquitoes is likely to affect other mosquito species that are
112 pollinators or otherwise known to be beneficial to ecosystem processes;
- 113 (g) Whether the LM mosquitoes are likely to have an adverse effect on other interacting organisms,
114 e.g. predators of mosquitoes;

- 115 (h) Whether species replacement by other disease vector species may occur, and if so, whether it
116 can result in an increased incidence of the target disease or new diseases in humans or animals.

117 **Gene Flow**

118 (See steps 2 and 3 of the Roadmap for Risk Assessment of LMOs)

119 *Rationale:*

120 With regard to the biosafety of LM mosquitoes, gene flow refers to the transfer of transgenes⁴⁹ or genetic
121 elements from the LM mosquitoes to non-LM mosquitoes. It can occur via cross-fertilisation or other
122 movement of the transgenes or genetic elements. Various factors may influence gene flow and any
123 associated adverse effects, such as, the strategy, the transgenes, the gene drive system⁵⁰ and the stability
124 of the trait(s) carried by the mosquito over generations, as well as the receiving environment, etc.

125 *Gene flow through cross-fertilization:* Some LM mosquitoes are being developed to spread the
126 introduced trait rapidly through the target mosquito population. For instance, when introduced into
127 *Anopheles gambiae*, the trait may be expected to spread throughout the *A. gambiae* species complex.
128 Other LM mosquito technologies are designed to be self-limiting and, in such cases, spread of the
129 transgenes or genetic elements in the target mosquito population is not intended or expected. For the self-
130 limiting technologies, the potential for an unexpected spread of the introduced trait should be considered
131 by focusing on the assumption that any management strategy to limit the spread could fail. Gene flow
132 between different species should be considered for all of the LM mosquito technologies in spite of the
133 fact that mosquitoes, like other insects, typically have strong reproductive isolating mechanisms that will
134 not allow interspecific gene flow. Identifying the key reproductive isolating mechanisms and possible
135 conditions that could lead to the breakdown of such mechanisms is of particular importance in the risk
136 assessment of LM mosquitoes with this trait. In addition, the fitness conferred by the introduced trait and
137 the population size and frequency of the introduction of the LM mosquito into the environment will also
138 determine the likelihood and rate of spread of the transgenes or genetic elements.

139 *Horizontal gene flow:* For the purpose of this document, “horizontal gene flow”, is the movement of
140 genetic information from one organism to another through means other than sexual transmission. Gene
141 drive systems for moving genes into wild populations may be the initial focus of the risk assessment. The
142 risk of horizontal gene flow in LM mosquitoes that do not contain a gene drive system is likely to be
143 smaller but should nevertheless be assessed on a case-by-case basis.

144 *Persistence of the transgene in the environment.* Some of the transgenes in LM mosquitoes are designed
145 not to persist whereas others are expected to spread rapidly and/or persist through wild populations. In
146 cases where the LM mosquitoes have been found through the risk assessment process to have the
147 potential to cause adverse effects to the biological diversity, taking also into account human health,
148 methods to reduce the persistence of the transgene in the environment needs to be considered

⁴⁹ For the purpose of this document, a transgene is a nucleic acid sequence in an LMO that results from the application of modern biotechnology as described in Article 3 (i) a of the Protocol.

⁵⁰ Gene drive systems are methods of effectively introducing the desired gene into a mosquito population (Selfish DNA versus Vector-Borne Disease, Environmental Health Perspectives (2008) 116 - <http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2235231/pdf/ehp0116-a00066.pdf>).

149 *Points to consider:*

- 150 (a) Whether LM mosquitoes have the potential to transfer the modified traits to wild mosquito
151 populations (when it is not an intended strategy) and/or to non-related organisms, and if so, the
152 occurrence of any potential undesirable consequences;
- 153 (b) Whether the LM mosquitoes have the potential to induce undesirable characteristics, functions,
154 or behaviour within the target mosquito species, other wild related species or non-related
155 organisms;
- 156 (c) Any undesirable consequence should the transgene persist in the environment.

157 **Evolutionary responses (especially in target mosquito vectors or pathogens of humans and**
158 **animals)**

159 *(See step 1 of the Roadmap for Risk Assessment of LMOs)*

160 *Rationale:*

161 Any strong ecological effect also exerts an evolutionary selection pressure on the human and animal
162 pathogens and the mosquito vectors. The main evolutionary effects are those that could result in a
163 breakdown in the effectiveness of the technology and the resumption of previous disease levels. Some
164 LM mosquito strategies aim at modifying the mosquito vector's ability to transmit diseases through
165 changes in its physiological mechanisms. An evolutionary effect resulting in the development of
166 resistance to physiological mechanisms in the targeted pathogen might occur when modifying mosquito
167 vector competence. This might harm the effectiveness of the strategy used and result in a population of
168 pathogens that may be transmitted more easily by all types of vectors.

169 Other evolutionary effects could be hypothesized, including effects resulting from climate change, but
170 they would first require the occurrence of some adverse effect on a species, community or ecosystem
171 effect. Therefore, consideration of secondary evolutionary effects can be postponed until such effects are
172 identified and found to be significant.

173 *Points to consider:*

- 174 (a) Whether the target mosquito vector has the potential to evolve and avoid population
175 suppression, regain vector competence or acquire new or enhanced competence to another
176 disease agent, and if so, the occurrence of any possible undesirable consequences;
- 177 (b) Whether the trait has the potential to evolve and thus lose its effectiveness, or the pathogen to
178 evolve and overcome the limitation posed by the genetic modification, and if so, the occurrence
179 of any possible undesirable consequences.

180 **RISK-MANAGEMENT STRATEGIES**

181 *(See step 5 of the Roadmap for Risk Assessment of LMOs)*

182 Risk assessors may want to consider risk-management strategies such as the quality control of the
183 released LM mosquitoes and monitoring them and the environment for potential unintended adverse
184 effects. There should also be strategies in place for halting the release and application of mitigation
185 methods if an unanticipated effect occurs. Careful implementation of the technology including the
186 availability of mitigations measures (such as an alternative set of control measures should a problem
187 occur) and the integration of other population control methods should be considered. In some

188 circumstances methods to reduce the persistence of the transgene in the environment or to mitigate
189 adverse effects resulting from the expression of the transgene might be needed. Monitoring during and
190 after the environmental release of the LM mosquitoes so as to address prompt detection of unexpected
191 adverse effects may also be considered.

192 *Points to consider:*

193 (a) Availability of monitoring methods to:

194 (i) Measure the efficacy and effectiveness of LM mosquito technology;

195 (ii) Assess the potential evolutionary breakdown of the LM mosquito technology (monitoring
196 for transgene stability and proper function over time);

197 (iii) Determine the level to which the identified adverse effects may be realized, including
198 detection of unexpected and undesirable spread of the transgenic trait (monitor for
199 undesirable functions or behaviours within target species and other wild related species).

200 (b) Availability of mechanisms to recall the LM mosquitoes and transgenes in case they spread
201 unexpectedly (e.g. mass release of wild-type mosquitoes above a certain threshold, alternative
202 control methods including genetic control).

203 (c) Availability of methods for managing the dispersal of the LM mosquitoes and ensuring that they
204 do not establish themselves beyond the intended receiving environment (eg. vegetation-free
205 zones, traps, high threshold gene drive systems).

206 (d) Availability of methods to manage potential development of resistance, e.g. in the target vector
207 or pathogen.

208 **OTHER ISSUES**

209 There are other factors that may be taken into consideration in the decision for environmental releases of
210 LM mosquitoes which are not covered by Annex III of the Protocol. They encompass, *inter alia*, social,
211 economic, cultural and health issues associated with the application and acceptance of the technology.

212 **BIBLIOGRAPHIC REFERENCES**

213 See references relevant to the "[*Guidance Document on Risk Assessment of LM Mosquitoes*](#)".

*Annex IV***RECOMMENDATIONS TO THE CONFERENCE OF THE PARTIES SERVING AS THE MEETING OF THE PARTIES TO THE CARTAGENA PROTOCOL ON BIOSAFETY AT ITS FIFTH MEETING**

1. The Ad Hoc Technical Expert Group (AHTEG) on Risk Assessment and Risk Management took note of the deliberations under the Open-ended Online Expert Forum on Risk Assessment and Risk Management in particular about the need for further guidance on specific aspects of risk assessment and considered the existing guidance materials on risk assessment of living modified organisms.
2. The AHTEG recognized the importance of involving experts in the various scientific and technical fields relevant to risk assessment in any future activity taking into account the limited financial and human resources.
3. The following recommendations were made by the AHTEG:
 - (a) The document “Guidance on Risk Assessment of Living Modified Organisms” should be published and distributed, including an online version under the Biosafety Clearing-House (BCH), in all UN languages;
 - (b) The “Guidance on Risk Assessment of Living Modified Organisms” should be further tested for example during regional workshops including cooperation with existing initiatives for capacity-building and training, as appropriate;
 - (c) The “Guidance on Risk Assessment of Living Modified Organisms” should be revisited within two years and the need for an update of the list of background materials should be assessed within a year;
 - (d) Further development of guidance on risk assessment of living modified organisms should be considered. The topics identified and prioritized during the first meeting of the AHTEG as well as those mentioned at the second meeting could be the starting point for the further development of guidance on risk assessment (see list annexed hereto as annex V);
 - (e) A process should be established for the incorporation of background materials, available in the Biosafety Information Resources Centre of the Biosafety Clearing-House, that are relevant in the different sections of the “Guidance on Risk Assessment of Living Modified Organisms”. In order to assist this process, the Secretariat should be requested to revise the common format for submission of records to the Biosafety Information Resources Centre (BIRC) of the BCH with the view to identifying and including a mechanism to link BIRC records on risk assessment to specific sections of the guidance document;
 - (f) Recognizing that the exchange of information is a central element for identifying living modified organisms or specific traits that have been assessed as having the potential to cause adverse effects on the conservation and sustainable use of biological diversity taking also into account risks to human health, a process should be established by:
 - (i) Urging Parties and inviting non-Parties to submit relevant information to the BCH on experiences in conducting risk assessment with regard to this topic;

- (ii) Requesting the Secretariat to undertake a regular analysis of the information contained in the BCH within the context of this process and reporting to the COP-MOP for that purpose;
- (iii) Organizing workshops where the information submitted would be analyzed through a guided-process.

(g) The goals of the above recommendations (a) to (f) could be achieved by a combination of an extended Open-ended Online Expert Forum on Risk Assessment and Risk Management and an AHTEG on Risk Assessment and Risk Management, as well as a combination of online conferences, ad hoc discussion groups and face-to-face meetings with a view to:

- (i) Developing additional guidance documents on the basis of the “Guidance on Risk Assessment of Living Modified Organisms” on specific types of living modified organisms and traits;
- (ii) Reviewing the text of the “Guidance on Risk Assessment of Living Modified Organisms” and updating the lists of background materials;
- (iii) Incorporating background materials, available in the Biosafety Information Resources Centre of the Biosafety Clearing-House, that are relevant to the different sections of the “Guidance on Risk Assessment of Living Modified Organisms”;
- (iv) Analysing the results of the workshops on living modified organisms or specific traits that have been assessed as having the potential to cause adverse effects.

(h) Human and financial resource implications should be considered for the process set up to achieve the above goals.

Annex V

TOPICS FOR THE DEVELOPMENT OF GUIDANCE MATERIALS ON RISK ASSESSMENT

*Further topics indentified in the first meeting of the AHTEG as priorities for the development of guidance:*⁵¹

- Post-release monitoring and long-term effects of LMOs released into the environment;
- Risk assessment and risk management in specific receiving environments;
- Risk assessment of living modified microorganisms and viruses;
- Risk assessment of living modified pharmaplants;
- Risk assessment of living modified crops;
- Risk assessment of living modified trees;
- Risk assessment of living modified fish;
- Risk assessment living modified organisms for production of pharmaceutical and industrial products;
- “Co-existence” between LMOs and non-LMOs in the context of small scale farming;
- Risk assessment of living modified plants for biofuels;
- Risk assessment of living modified organisms produced through synthetic biology.

Further topics identified in the second meeting of the AHTEG as possible priorities for the development of guidance:

- Uncertainty analysis;
- Establishment of criteria for transparency and reproducibility of information;
- Interface between risk assessment and risk management;
- Environmental risk assessment and monitoring taking into account human health;
- Unintentional transboundary movements;
- Risk assessment and management of LMOs intended for introduction into unmanaged environments.

⁵¹ From annex II of the report of the first meeting of the Ad Hoc Technical Expert Group on Risk Assessment and Risk Management (UNEP/CBD/BS/COP-MOP/5/INF/13).