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Certified Reference Materials AOCS 0809-A2

Report of the certification process for

MON 87701

Soybean Certified Reference Material

Second Batch

OECD Unique ID MON-877Ø1-2

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Abstract

This report describes the preparation and certification of the soybean CRM AOCS 0809-A2 produced by AOCS Technical Services in 2019. The CRMs have been prepared according to ISO 17034:2016 and are intended to serve as control material for third party testing of soybean for transformation events. The soybean MON 87701 powder was provided by Bayer CropScience, St. Louis, MO. It was prepared by grinding the bulk seed at Bayer CropScience. The certified value of AOCS 0809-A2 was based on the purity of the bulk seed material and with 95% confidence, the true value is ≥ 968 g/kg. The powder was aliquoted and bottled in 27-mL glass headspace vials and sealed under a nitrogen gas environment at Illinois Crop Improvement Association. The presence of MON 87701 in AOCS 0809-A2 was verified using event-specific, qualitative PCR analysis by Eurofins-GeneScan, New Orleans, LA (an ISO 17025 accredited laboratory). CRM samples should be stored in a dry, sealed container at ambient or cooler conditions in the dark.

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Glossary

AOCS American Oil Chemists' Society

Conventional Crop Crop variety with no history of transgenic technology and is

produced through traditional plant-breeding techniques that rely on selecting and mating parent plants possessing promising traits and repeatedly selecting for superior

performance among their offspring

DNA Deoxyribonucleic Acid is the linear, double-helix

macromolecule that makes up the genetic material of most

organisms

Detection Limit Lowest level at which target DNA can be detected in a sample.

EC European Commission

Genome The full set of genes and associated DNA characteristic of an

organism

ISO International Organization for Standardization

GMO Organism that has had genetic sequences modified using

molecular-level techniques

PCR Polymerase Chain Reaction: technique used to determine

whether a sample of plant tissue contains a particular DNA sequence. PCR relies on primer sets that zero in on a

particular target DNA sequence and a special DNA-copying enzyme (DNA polymerase) that makes enough copies of the

target sequence for identification and measurement

Qualitative PCR PCR methods that determine the presence or absence of a

specific target DNA sequence at a particular level of detection

Report of Certification for 0809-A2 Page 6 of 12 ©AOCS, 2024 Quantitation Limit Lowest level at which the amount of target DNA sequence in

a sample can be reproducible.

Quantitative PCR PCR methods that estimate the relative amount of target DNA

sequence in a mixture of DNA molecules

Trait: MON 87701 Homozygous trait for insect-protected soybeans that

produces the Cry1Ac insecticidal crystal protein derived from

Bacillus thuringiensis (Bt) subsp. kurstaki. The Cry1Ac protein

provides protection from feeding damage caused by targeted

lepidopteran pests

Introduction

Plant genetic modification is an extension of traditional plant breeding. It allows plant breeders to develop crops with specific traits including insect, disease, and herbicide resistance; processing advantages; and nutritional enhancement. An important component for identifying these new traits is a Certified Reference Material created from leaf, seed, or grain containing the new trait as well as a CRM created from the conventionally bred matrix. The European Commission has mandated that from 18 April 2004, a method for detecting a new event derived from transgenic technology and Certified Reference Material must be available before the EC will consider authorizing acceptance of a new crop derived from transgenic technology. Several nations outside Europe also require grain and ingredients to be labeled above a threshold level before accepting a shipment.

To meet the above regulatory requirements for GMO determination, AOCS 0809-A2 was manufactured from soybean according to ISO 17034:2016 and in accordance with EC No 1829/2003, EC No 641/2004 and EC No 619/2011. This CRM is available from AOCS.

Material Processing

MON 87701 soybean seeds used to prepare AOCS 0809-A2 were homozygous resulting from several cycles of self-pollination. Bayer CropScience milled ~7 kg of MON 87701 soybean seed. All of the seed powder was passed through a 710 μM mesh sieve. The seed powder was delivered to AOCS who contracted Illinois Crop Improvement Association for packaging the samples. The powder was aliquoted and bottled in 27-mL glass headspace vials and sealed under a nitrogen gas environment.

Trait Verification to Certify Presence of MON 87701

The presence of the MON 87701 trait was assessed on 10 random vials of AOCS 0809-A2. AOCS used the Random Number Generator function of Microsoft Excel to select samples for verification of trait presence. Sample numbers that were randomly

selected were sent to Eurofins-GeneScan, New Orleans, LA (an ISO 17025 accredited laboratory) for event-specific, qualitative PCR analysis to verify the presence and homogeneity of MON 87701 in the samples (Table 1).

Table 1. Trait verification and testing on AOCS 0809-A2 MON 87701 soybean performed by Eurofins-GeneScan, New Orleans, LA (an ISO 17025 accredited laboratory).

AOCS 0809-A2 Sample	Trait MON 87701 Presence
Sample # 0005	Positive
Sample # 0190	Positive
Sample # 0031	Positive
Sample # 0011	Positive
Sample # 0174	Positive
Sample # 0180	Positive
Sample # 0015	Positive
Sample # 0128	Positive
Sample # 0161	Positive
Sample # 0108	Positive

Certified Value and Measurement Uncertainty

The genetic purity of the seed lot used to produce AOCS 0809-A2 was assessed by Bayer CropScience. A total of 720 soybean seeds were subjected to individual seed testing for the presence of MON 87701 by qualitative event-specific PCR. 714 of the 720 seeds tested positive for the presence of MON 87701.

Purity estimation was calculated using SeedCalc8 (Remund *et al.*, 2008) and corresponded to the lower bound of the true % purity. The % purity in the sample was 99.2% when 720 seeds were tested. Using a 95% confidence level, the true % purity of the MON 87701 seed lot was 96.8%. Consequently, with 95% confidence, the true value is \geq 968 g/kg.

The Measurement Uncertainty was based on the lower bound of the true % purity. The measurement uncertainty is the expanded uncertainty with a coverage factor of 2 and confidence level of 95%. It is obtained by combining the uncertainties from the purity assessment $(u_{char,rel})$, the homogeneity assessment $(u_{bb,rel})$, the transport stability assessment $(u_{sts,rel})$ and the long-term stability assessment $(u_{lts,rel})$:

$$u_{CRM,rel} = \sqrt{u_{char,rel}^2 + u_{bb,rel}^2 + u_{sts,rel}^2 + u_{lts,rel}^2}$$

$$U_{CRM} = 2 \times u_{CRM,rel} \times 1000 \ g/kg$$

When using an asymmetric uncertainty, the reported measurement uncertainty is truncated on the right side such that the value does not exceed 1000 g/kg. Consequently, the expanded measurement uncertainty for AOCS 0809-A2 is +8 g/kg, -24 g/kg.

Homogeneity

The homogeneity of AOCS 0809-A2 is related to the purity of the seeds. 714 out of 720 seeds tested positive for the MON 87701 soybean event by event-specific PCR. Based on the sample purity of 99.17%, as determined using SeedCalc8, the batch was considered to be homogenous.

In addition, the homogeneity of the MON 87701 trait was confirmed when 10 random vials of AOCS 0809-A2 were selected and were sent to Eurofins-GeneScan, New Orleans, LA (an ISO 17025 accredited laboratory) for event-specific, qualitative PCR analysis to verify the presence of MON 87701 in the samples (See Trait Verification section and Table 1).

Stability

Time, temperature and light are regarded as the most relevant influences on the stability of CRM (Linsinger, et al., 2001). The influence of light is mitigated by shipping and storing the vials in boxes, thus minimizing the possibility of degradation due to light. The influence of temperature is mitigated by storing the vials in a temperature-controlled room, and shipping vials at ambient temperature.

The effect of temperature and time are investigated.

stability is 1.0%.

A transport (short-term) stability study is conducted to assess the stability of maize CRM during transport. The temperature and time conditions in the study cover the typical conditions and the not so rare situations. The outcome of the study is considered transferable to other CRMs of similar property. Samples were subject to 3 different temperatures (4 °C (fridge), 25 °C (ambient), 60 °C (oven)) for 4 different durations (0, 1, 2, and 4 weeks). The study concluded that samples are stable at 4 °C (fridge) and 25 °C (ambient) for 4 weeks. The estimated uncertainty contribution from transport (short-term)

A long-term stability study is conducted to assess the stability of maize CRM during storage. Samples are stored at 25 °C (ambient) and the stability of the sample is monitored as long as the samples is available. The storage temperate studied is 25 °C and the length of time to be studied is 10 years. The outcome of the study is considered transferable to other CRMs of similar property. In the initial 1-year stability study, samples were subject the storage condition for 4 different durations (0, 1, 3, 6 and 12 months). The study concluded that samples are stable at 25 °C (ambient) for 12 months. The estimated uncertainty contribution from long-term stability is 0.42%.

Stability of these CRMs has been listed as 1 year from the certification date. The materials were processed and are stored at ambient temperature, under nitrogen gas, in 27 -mL glass headspace vials. These materials are expected to be stable for longer than the estimated expiration date. The stability of the powder material will be reevaluated at time of expiration. If the samples still test positive for the presence of the intended trait, the certificates will be extended.

References

Eurofins-GeneScan; 2219 Lakeshore Drive, Suite 400, New Orleans, LA 70122; Telephone: +1 504 297 4330 Toll Free: +1 866 535 2730 Fax: +1 504 297 4335 https://www.eurofinsus.com/food-testing/testing-services/gmo/

Illinois Crop Improvement Association, 3105 Research Road, Champaign, IL 61826; Telephone: +1 217 359 4053 Fax: +1 217 359 4075; http://www.ilcrop.com/index.htm

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