
**Živila - Določevanje alergenov v živilih z molekularno biološkimi metodami - 7.
del:Arašidi (*Arachis hypogaea*) - Kvalitativno določanje specifičnega zaporedja
DNK v živilih s PCR v realnem času**

Foodstuffs - Detection of food allergens by molecular biological methods - Part 7: Peanut (*Arachis hypogaea*) - Qualitative detection of a specific DNA sequence in food by real-time PCR

Lebensmittel - Nachweis von Lebensmittelallergenen mit molekularbiologischen Verfahren - Teil 7: Erdnuss (*Arachis hypogaea*) - Qualitativer Nachweis einer spezifischen DNA-Sequenz in Lebensmitteln mittels Real-time PCR

Produits alimentaires - Détection des allergènes alimentaires par des méthodes d'analyse de biologie moléculaire - Partie 7 : Arachide (*Arachis hypogaea*) - Détection qualitative d'une séquence d'ADN spécifique dans des produits alimentaires, par PCR en temps réel

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Qualitative detection of a specific DNA sequence in food by
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Lebensmittel - Nachweis von Lebensmittelallergenen
mit molekularbiologischen Verfahren - Teil 7: Erdnuss
(*Arachis hypogaea*) - Qualitativer Nachweis einer
spezifischen DNA-Sequenz in Lebensmitteln mittels
Real-time PCR

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CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels

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