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## Multi-faceted Migration in Food contact Polyethylene-based Nanocomposite

2 packaging

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- 8 ABSTRACT:
- 9 The flourishing market of nanocomposite food packaging has raised concerns about the safety of 10 these materials. While several works on this issue have been published in recent years, they main 11 focus in these studies was found to be on the possible migration of the nanoparticle its 12 constituents. However, thorough safety evaluation of these materials would not be realistic until 13 the nano-packaging system would be regarded as a whole with all of its components and the 14 interactions of all these components. This matter is specifically crucial in terms of the interaction 15 of nanoparticles with the non-nano additives which are added during the packaging processing. 16 As the toxicity of these processing additives is no less than the nanoparticles, the possible impact 17 of the nanoparticles on the transfer properties of these substances could play a decisive role on 18 the risk assessments of the nanocomposite for food application. This study is an attempt through 19 a thorough analysis of nanocomposite risks in terms of the interactions of components and the 20 resulting effects on the release of nanocomposite substances. In this regard a model 21 nanocomposite of LLDPE and nanoclay which is also comprised of intercalants and some 22 selective additives were put in contact with various food simulants were considered and the

global, specific and elemental exposure to the substances were extensively evaluated. The results
 are believed to provide more tenable judgements about the safety of polymer nanocomposites.
 KEYWORDS: Clay Polymer Nanocomposite, Multi-dimensional Migration, Food contact
 materials regulations

#### 1. Introduction

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To comply with shelf-life requirements, polymer nanocomposites (PNC) have emerged in the past decade as a new class of food packaging materials which provides several advantages, such as enhanced mechanical, thermal and barrier properties of conventional plastic packaging or improved function of active (antimicrobial) and intelligent materials (Farhoodi, 2016; Gontard et al., 2017). Particular interest is that such improvement in nanocomposite properties are achieved with very low loading levels (typically up to 5% by weight) of nanoparticles while these properties are not commonly attained with less than 25-40 wt% loading in the microcomposites (Chaudhry et al., 2017). However, besides the advantages of nanocomposite application, their safety issue should be taken into consideration especially for food packaging systems, because of the potential adverse effects of nanoparticles on human health. Nevertheless, the "safety aspects" in most of recent researches are considered in respect to the potential migration of nanoparticles from different polymers to several foods and in diverse conditions of contact time/temperature (Metak et al., 2015) or to the toxicity of these nanoparticles for human health and environment (Gaillet and Rouanet, 2015; Sajid et al., 2015). In viewpoint of the regulations, scarce rules could be found due to the lack of information on the risk assessment of nanoparticles, and even among these regulations there is not a consensual management on the usage of nanoclay in Food Contact Materials (FCM). According to the Food and Drug Administration (FDA), montmorilonite (Mt) is considered as GRAS (Generally Recognized As Safe, under the section 21CFR184.1155) and can be consequently used in food packaging formulation (Molinaro et al., 2013). This statement is based on the in vitro studies indicating the cytotoxic effects of clay minerals in case of exposure to a high dose (e.g. 53 thousands of ppm), which is unlikely to happen for human being (Li et al., 2010; Baek et al., 54 2012). 55 In contrast, the European Food Safety Authority (EFSA) in a recent guidance on the risk 56 assessment of nanotechnologies in the food and feed chain (Hardy et al., 2018) considered that 57 the scientific knowledge remains too limited today to define a threshold for the migration of nanomaterials from FCM. As consequence, EFSA recommend that "an argument for safety may 58 59 be made on a case-by-case basis if migration of a nanomaterial in particulate form is only in 60 trace amounts". 61 Among the engineered nanomaterials used in packaging placed on the market, Mt is already 62 widely employed to improve the barrier properties (oxygen and carbon dioxide permeability) of 63 conventional materials. Mt is the main component of bentonite which is included in the list of authorized substance (No 393) of the Regulation (EU) 10/2011 (European Commission, 2011) 64 65 without restriction. Recently the EFSA Panel on FCMs, enzymes, flavorings and processing aids 66 (CEF) has amended a scientific opinion on the use of montmorillonite clay in FCMs which 67 concludes that inclusion of nanoclay up to 12% w/w do not represent a health risk because of the absence of potential migration (EFSA Panel on Food Contact Materials, Enzymes, 2016). In 68 69 spite of the published guidance on the safety evaluations of nanomaterials, the scientific 70 committee on emerging and newly identified health risks reported these procedures could be 71 expected to be remained under development until there would be sufficient information available 72 to characterize the possible harmful effects on human and the environment (Williams, 2006; 73 Wyser et al., 2016). 74 The toxicity and exposure to the nanoparticles are crucial factors in evaluating the risks related to 75 their food packaging application, by limiting the studies to these two aspects, several other

sources of risks being still ignored. In order to better apprehend the risks of nanocomposites for such application, an exhaustive investigation must include additional components and their interactions. In a simplistic view, the nanocomposite packaging in contact with food could be considered to comprise three main component systems i.e. polymer, nanoparticles and food. However, several other additives (plasticizers, anti-oxidants, UV stabilizers, anti-static agents) are usually added to the plastic-based material during the packaging processing to further enhance their properties. Moreover, for most of inorganic nano-clays such as Mt, some other chemicals should be incorporated in the nanocomposite structure in order to disentangle the nanoparticle aggregates and consequently induce their exfoliation on one hand, and to enhance their compatibility with the base polymer in regards to the polarity on the other hand. Therefore, in an ordinary nano-packaging, a pentagon of interactions is involved between polymer, nanoparticle, food, processing additives and intercalants (the headspace volume to be considered as a possible sixth contributing part). It is worth noting that processing additives and intercalants are not considered in the same category because the type and quantity of the latter have direct and decisive effect on the efficiency of the nanocomposite as a "nano"-structure. In polymers with low polarity (such as polyolefins) the chemical modification of the polymer using a compatibilizer (adhesion promoter) such as maleic anhydride-grafted polyethylene (MA-g-PE) is necessary to promote greater interaction of the matrix with the filler and facilitate the formation of exfoliated nanocomposite structures. Although increasing the quantity of compatibilizer as well as modifier to a certain level facilitates the dispersion of nano-clay through the base polymer, these components should be examined for their migration as it should not surpass the authorized level.

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In addition to the migration of the nanoparticles and the intercalating agents, and like any other FCM, safety assessment of nanocomposite packaging must comply with the general requirement of the EU regulation that define overall and specific migration limits (European Commission, 2011). Overall migration (OM), the total mass released by the packaging material is regarded as an indicator of the food contact material inertness which sets an upper limit of 10 mg for all transferred substituents per dm<sup>2</sup> (or 60 mg/kg food) of the food contact article. On the other hand, based on the toxicological information of the molecules, the specific migration limits (SML) were set out for a single component to ensure the safety of the final package. Preliminary step in evaluating the safety of the nanocomposite is to determine its overall migration which takes into account the total release of nanoparticle (or its components), modifier, compatibilizer, residual monomers, processing additives and the non-intentionally added substances (NIAS) which may be present in the nanocomposite in the form of impurities in the additives, monomers and other starting substances which are used to manufacture the nanocomposite. The second step concerns the assessment of nanoparticle migration which has attracted the most attention in the recent years and could be considered from three different perspectives: Regulations, theory and experiments. So far, no evidence was found in the literature to indicate that nanoclay is likely to cause adverse effects on health when used in the food packaging (Piperigkou et al., 2016). However, while several studies provide some information about the migration of nanoclay/nanoclay constituents, the effects of nanoclay incorporation on the potential migration of chemical constituents of plastics (monomers and processing additives) were not taken into consideration.

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The third part of the nanocomposite safety evaluation is related to the migration of the compatibilizer and the modifier which is of significance due to the two factors related to the compliance with SML and alteration of the nature of the nanocomposite. In other words, not only the migration limits of these components should be less than the values which are set out in Commission Regulation (EU) No 10/2011, but also the effect of their release on the nature of the nanocomposite (in terms of polarity) and the consequences on the adsorption/desorption of other additives should be evaluated.

In the last step, the effect of the nanoparticle and its accompanying components (modifier and compatibilizer) on the migration of several additives with various size, functional groups, polarities and volatilities was assessed according to a specifically designed challenge test which was originally recommended by EFSA to determine the decontamination efficiency of the recycling process for recycled polymer packaging (EFSA, 2011).

As a consequence of the multiple factors involved in the safety evaluation of nanocomposite packaging in terms of the exposure to various migrants, this study attempts to draw a broader perspective of the possible interactions between the nanocomposite components and the release

#### 2. Experimental

2.1. Chemicals and food simulants

Linear low-density polyethylene (LLDPE, LL 1002YB melt flow index 2.0 g/10 min, density 0.918 g/cm3) was supplied by Exxon Mobil Chemical. Cloisite 20 (C20), a Bis(hydrogenated tallow alkyl)dimethyl ammonium bentonite salt (Bulk density: 350 kg/m3; Density (20 °C): 1.80

of diverse substances in contact with different type of food simulants.

142 g/cm3; Particle Size, D50: < 10 μm; Lamellar spacing (XRD, d001): 2.7 nm) was provided by 143 BYK Additives & Instruments. 144 Fusabond E226 (supplied by DuPont<sup>TM</sup>) was chosen as the compatibilizing agent between 145 nanoclay and the matrix which is a maleic anhydride modified polyethylene (MA-g-PE) with 146 melt flow index of 1.75 g/10 min and density of 0.93 g/cm3. 147 The chemical additives comprising of three solid surrogates (biphenyl 99.5%, benzophenone 148 99% and methyl stearate 99%), five liquid contaminants (toluene 99.8%, chlorobenzene 99.8%, 149 methyl salicylate ≥ 99%, phenyl cyclohexane ≥ 97% and DEHA ≥ 99%) as well as Arquad® 150 2HT-75 (Di-hydrogenated tallow di-methylammonium chloride) and Maleic anhydride 99% 151 were all provided by Sigma-Aldrich. 152 The migration of chemicals was evaluated using 5 food simulating liquids according the EU regulation 10/2011. The selected food simulants encompass 3w/v% aqueous acetic acid to 153 154 represent acidic food, ethanol 10% to simulate aqueous food, ethanol 50% for alcoholic foods 155 (with alcohol content of greater than 20%) as well as the dairy products and ethanol 95% and

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2.2. Polymer Nanocomposite (PNC) processing

iso-octane which are both representing fatty foods.

The nanocomposite was synthesized with LLDPE as the base matrix, 5 wt% of C20 and 15 wt% of MA-g-PE by melt intercalation method using a co-rotating twin screw extruder (Thermo Scientific<sup>TM</sup> EuroLab 16) with a L/D ratio of 40 and a screw diameter of 16mm at screw speed of 200 rpm and feed rate of 1.0 kg/h.

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#### 164 2.3 Impregnation of PNC

The virgin polymer and nanocomposite were contaminated by a mixture of additive with theoretical additive amount of 500-1000 ppm, which is within the range recommended by EFSA (EFSA, 2011), The Impregnation of PNC was performed with 80 µl of additive cocktail mixed with 25 g of the polymer pellets. The bottles were sealed and stored at 40 °C under rotary agitation. After 1 week, the contents were rinsed with distilled water and the contaminated pellets were divided and subjected to the four chosen food simulants as well as the pure dichloromethane for extraction to obtain initial concentration ( $C_0$ ). The polymer samples were also extracted with pure dichloromethane to evaluate the initial quantity of additives (M<sub>0</sub>) before the migration process by gas chromatography (GC-FID, Agilent 7890A gas chromatograph equipped with a HP-5 capillary column of 32mm × 30m ×

## 2.4 Overall Migration Test

 $0.25 \, \mu m$ ).

Overall migration (OM) level which represents the total amount of non-volatile substances transferred from the food-contact material to the food (EU 10/2011) is determined by a gravimetric method after the total immersion of neat samples in the selected food simulants based on the standardized testing conditions set out for long term storage (40°C for 10 days) in EU 10/2011.

In this regard, the neat samples were cut in disks of 3 cm diameter and then kept in a vacuum oven at 50°C for 2 days. Respecting a surface/volume ratio of 6 dm²/L of food simulant, the polymer disks of a thickness of 177.4±16.1 and 143.3±8.3 µm for LLDPE and PNC respectively with the total area of 60 cm² were weighed using a microbalance (0.1 mg resolution) prior to the

migration test and then placed in a 100 ml migration cell (DURAN® screw thread tubes) with PTFE-sealed caps. Such a sample geometry makes it possible to neglect any migration linked to a possible edge effect. Glass beads were used as spacers between the disks to ensure a proper contact with the simulant. Before filling, the migration cells were blown out with nitrogen to prevent any dust contamination. The cells were then filled with the respective simulants, sealed and stored in a 40°C oven. Each sample/simulant combination was prepared in triplicate for the migration test.

After 10 days of contact, the bottles were ultrasonicated for 2 min to detach any potential particles from the sample surface and the glass beads. The samples then removed and placed in a

vacuum oven at 50°C for 3 days. After evaporation of the simulant, the specimens were weighed

again and eventually overall migration was calculated in milligrams per square decimeter of the

#### 2.5 Specific Migration Test

surface area of sample.

Specific migration test was performed according to recommendation described in EU 10/2011 under standard conditions identical to those described for the global migration test. The European legislation has published a list of substances allowed in food contact articles for which the specific migration limits are assigned based on the toxicological evaluation of the substance. These limits which are expressed in mg substance per kg food are 30mg/kg under the FCM Substance No. 234 and No. 15 for maleic anhydride and quaternary ammonium salt, respectively (European Commission, 2011). Therefore, the migration of these two components were evaluated by the same experiment as for the overall migration. By the end of the contact time,

the simulants were collected to be analyzed by HPLC for maleic anhydride and LC-MS/MS for di-hydrogenated tallow di-methylammonium chloride.

Except for the iso-octane, other simulants were directly injected into the HPLC and LC-MS/MS system. However, iso-octane needed additional sample preparation. In this regard, the simulant was evaporated to dryness by a nitrogen stream. The residue was then re-dispersed in ethanol and was severely agitated on a vortex for 30 s.

- 2.5.1 Liquid chromatography tandem-mass spectrometry (LC-MS/MS)
- LC-MS/MS analysis was performed on a Synapt G2-S (Waters) spectrometer equipped with a positive electrospray ionization (ESI<sup>+</sup>) source. The capillary voltage was set to 3.5 kV and the sampling cone voltage was set to 60 V. Compounds were separated using a gradient of methanol (solvent B) in 0.1% formic acid in water (solvent A) over 15 minutes. The gradient program was set as follows: 0-2 min, 20% solvent B; 2-3 min, 20-80% solvent B; 3-5 min, 80-95% solvent B; 5-13 min, 95% solvent B; and 13-15 min, 20% solvent B. Standard solutions of Arquad® 2HT-75 in ethanol, with concentrations ranging from 0.1 to 10 mg/L, were used to establish the external calibration curves based on the C<sub>18</sub>C<sub>18</sub> peak areas.

- 2.5.2 High Performance Liquid Chromatography (HPLC)
- Maleic anhydride concentration in simulants was analyzed by Alliance HPLC with a reversed phase Alltima C18 column (250 mm ×2.1 mm, 5-μm packing) protected with a guard column of Alltima C18 (7.5mm × 2.1 mm, 5-μm packing). Compounds were separated using isocratic elution with acetonitrile/water/phosphoric acid 30:70:0.1 (v/v/v). Elution flow rate was maintained at 0.2 mL/min and UV detection was recorded at 240 for ethanol 95% and pure

ethanol and 254 nm for other simulants based on the maximum absorption. External calibration was established in each simulant with concentrations of 0.1 to 100 mg L<sup>-1</sup>.

2.6 Inductively Coupled Plasma-Mass Spectrometry (ICP-MS)

Migration of nanoclay elements from the nanocomposite to the simulants was investigated by ICP-MS method. Elemental analysis measurements for Al, Si and Mg were carried out with a quadrupole ICP-MS instrument (iCAP Q, Thermo Scientific), equipped with a concentric nebulizer and a quartz cyclonic spray chamber connected to the ICP-torch for sample introduction. Before instrumental analysis, an acid digestion procedure is applied on the samples. After putting the neat samples in contact with the simulants for 10 days at 40°C, the samples were separated from the liquid and the simulants were evaporated under a nitrogen stream and re-dissolved in HNO<sub>3</sub> 1%. Blanks of the food simulants were also undergone the same procedure. All migration solutions were prepared in triplicate

Simulants were examined for the presence of (nano)particles as well as their elemental composition by SEM/EDX. In this regard, the simulants (after 10 days of contact with NC) were filtered through a Whatman Anodisc membranes with 0.2 µm pore size and 47 mm diameter. The filtrate liquid then passed through the Anodisc membranes with 0.02 µm pore size. The membranes were put in petri dishes to be dried in room temperature for 48 hours and covered with perforated aluminum foil to avoid the absorption of dust. After simulant evaporation, SEM/EDX analysis was carried out on the membranes using the EDAX attachment (Oxford Instruments) of the Hitachi S-4500 I SEM (Hitachi High Technologies America, Inc.,

2.7 Scanning Electron Microscopy with Energy Dispersive X-ray Spectroscopy (SEM/EDX)

Schaumburg, USA).

#### 3. Results and discussion

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258 3.1 Overall migration 259 Determining the overall migration of all components from the packaging is a key factor which is 260 typically conducted by the manufacturer or food packager to ensure the compliance with 261 European regulatory limit which is 10 mg/dm<sup>2</sup> on a contact area basis or 60 mg/kg in the 262 simulant or food. The global migration values for pure polymer and the PNC are reported in 263 Figure 1. 264 The total migration measured from PNC is higher than the corresponding values in pure 265 polymer, however, in viewpoint of the FCM safety, it is out of consideration as far as the values 266 are below the regulated limit. Although the OM value seems to transgress the limit in case of the 267 PNC contact with iso-octane, it should be pointed out that the contact condition applied in this study for iso-octane is 10 days at 40°C for the sake of the comparison with other simulants and 268 269 consequently higher than the recommended conditions of the European commission (2 days at 270 20°C) for checking the compliances with FCM regulations. Moreover, iso-octane represents a 271 very extreme condition as a food simulant due to the strong interaction between iso-octane and 272 the polymer. It was reported in the literature that even the contact duration of 2 hours at 40 °C 273 with this simulant results in a far greater migration than the 10 days of contact with olive oil at 274 the same temperature (Baner et al., 1992). The emphasized OM value in this simulant is largely 275 due to the iso-octane migration front which is penetrating into the polymer and is highly 276 depended on the nature of the polymer as well as the applied contact time. 277 Though the compliancy of overall migration in nanocomposites with the regulatory limits 278 mitigates the safety concerns about the application of these materials, it is of interest to recognize

the reasons for the elevated OM of nanocomposites in comparison to the pure PE. The mismatch

of the overall migration values with the sum of specific migration of nanoclay elements, modifier and compatibilizer, implies that the increased migration in nanocomposite can be explained by various comconmitting effects including (1) migration of oligomeric compounds from the partial degradation of LLDPE, (2) migration of additives (antioxidant type in particular) initially introduced into the commercial polymer fraction or (3) migration of impurities present in the sample of Cloisite 20A.

In addition and as evidenced by literature, the nanocomposite structure is prone to be affected by the stimulated thermo-oxidative degradation of polymer due to the nanoclay incorporation during melt-blending process. The rate of thermo-oxidative degradation was found to be increased by the oxygen-scavenging effect due to the char formation in the clay surface as well as the catalytic activity of the acidic active sites created by the Hoffman decomposition of the alkyl quaternary ammonium ions (Annamalai and J. Martin, 2014). During thermal degradation proceeding according to the Hofmann degradation mechanism the ammonium cation loses an olefin and an amine and leaves an acid proton on the surface of the Mt. This acid site on the surface of Mt has a catalytic effect during the initial stages of decomposition of organic material within the organomodified nanoclay (Leszczyńska et al., 2007).

It should be noted that although the synthesized nanocomposite in this work was exposed to 160-180°C in the extruder, which is just before the onset decomposition temperature of the alkyl quaternary ammonium ions (Xie et al., 2001), the masterbatch method of synthesis could trigger more degradation by heat dissipation due to the combined effects of higher filler content and higher melt viscosity (Shah and Paul, 2004). Moreover, the heat barrier effect of nanoclay could provide superheated conditions inside the polymer melt leading to extensive random scission of polymer chain and evolution of numerous chemical species which, trapped between clay layers,

have more opportunity to undergo secondary reactions. As a result, some degradation pathways could be promoted leading to enhanced charring (Leszczyńska et al., 2007).

Another factor contributing to the polymer degradation is the presence of compatibilizer. Melt processing of PE with PE-g-MA is considered to ease PE chain scission (Esteki et al., 2013) or initiating some other side reactions (crosslinking) which would cause some changes in the molecular structure of the polymer (Rzayev, 2011).

The presence of maleic anhydride as a compatibilizer between nanoclay and LLDPE lowers the

## 3.2 Migration of nanoclay modifier and compatibilizer

surface energy of nanoclay and improves the wetting characteristics of the polymer which consequently leads to a better dispersion of nanoclay in the polymer. However, in European legislation, a group restriction was specified for this substance which indicates that the total specific migration limit (SML<sub>(T)</sub>) for the sum of substances applicable to this group should be less than 30 mg/kg (European Commission, 2011). As it is demonstrated in Table 1, the migration of the compatibilizer is far below the specific limitation for this substance.

The migration process can be described as the result of the diffusion of chemical additives through the polymers and the desorption of the diffused molecules from the polymer surface to the food or food simulants. In addition to the transport properties of a component (*i.e* diffusivity and freedom of movement which could be restricted by obstruction effect of the nanoparticles), the migration of organic molecules such as intercalants and compatibilizer, additionally depends on their respective affinity for both the food simulants and PNC. The polarity of diffusing molecules therefore appears as a determining factor of food contamination.

The results reported in Table 2 demonstrated a main influence of the nature of food simulant with higher migration values observed for fatty food simulant. Regarding the migration of maleic anhydride which exhibits a hydrophilic character, no tendency towards iso-octane was noted due to the difference in nature in terms of their polarity. The migration of maleic anhydride proved to be dramatically enhanced in ethanol 95% that can be explained by a modification of polymeric structure due to the sorption of ethanol by the LLDPE-based material and subsequent increase of substance diffusivity. Nanoclay modifiers such as quaternary alkyl ammonium is primarily used to increase the nanoclay interlayer space, however, once the nanoclay were embedded into the polymer, the modifier may be released from the nanoclay surface into the polymer matrix during the polymer processing or when the nanoclay was in contact with the solvent due to the solvent penetration into the polymer matrix. This part of modifier is considered as "free" and its release may follow the diffusion behavior of small molecules within the polymer matrix due to the presence of free volume and polymer chain relaxation (Yining et al., 2014). The migration of modifier from the PNC is demonstrated in Table 2 for all simulants. As mentioned above, it could be inferred that in contact with iso-octane and ethanol 95%, the simulant easily penetrates the polymer, swells the matrix and interacts with the modifier, which leads to the higher release of modifier in comparison to other simulants. However, the results of this study is not in accordance with the EFSA opinion on modified Mt, in which migration of dimethyl dialkyl (C16-C18) ammonium compounds from polyolefins was not detected in ethanol 95% with a LOD of 6 µg/kg (EFSA Panel on Food Contact Materials, Enzymes, 2016). Nevertheless, even in the condition of this study, the migration of the modifier

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is less than the 30 mg/kg which is regulated under FCM No. 15 for "alkyl, linear with even number of carbon atoms (C12-C20) dimethylamines" (European Commission, 2011).

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3.3 Migration of Nanoclay or their elementary building blocks

Study of the transport of engineered nanoparticles (ENM) within nanocomposite and its release into different food is crucial in the safety assessment of the PNC. In addition to a possible contamination due to the mechanical erosion of the PNC, ENM release phenomena during the life cycle of nanocomposite could be taken place by diffusion, desorption and dissolution that could be accelerated by the degradation of the polymeric matrix. Although multiple processes for nanomaterial release may simultaneously occur, the main release process depends on where the nanomaterial is located, the extent to which it interacts with the surrounding media, its ability to migrate through the host matrix material (size and steric hindrance), and whether the particle remains an ENM or is transformed into ionic form (Noonan et al., 2014). The nanoclay used in this study is montmorillonite (i.e. bentonite), which is a natural 2:1 layered phyllosilicate (de Abreu et al., 2009; Hannon et al., 2015) exhibiting an average thickness of ~1nm and average lateral dimensions ranging from a few tens of nm to several µm. Each platelet contains a layer of aluminum or magnesium hydroxide octahedral sandwiched between two layers of silicon oxide tetrahedral, with the ideal formula (full unit cell) of M<sup>+</sup><sub>0.66</sub>Si<sub>8</sub>Al<sub>3.34</sub>Mg<sub>0.66</sub>O<sub>20</sub>(OH)<sub>4</sub> in which M<sup>+</sup> represents the cation charges in the interlayer (Uddin, 2018). As Si, Al and Mg are the 3 main elements of the nanoclay, the migration of these elements into the food simulants are measured by ICP-MS and represented in Figure 2.

Considering the safety limits for the migration of metallic element and in spite of the welldeveloped knowledge on their toxicity and assignment of the provisional tolerable intake (1 mg/kg bw/week (Panel on Food Additives, Flavourings, 2016) & 250 mg/person/day (European Food Safety Authority, 2006) for Al and Mg respectively), the specific requirements for metals in food contact materials specified in Annex II of the Regulation EU/10/2011 does not include any specific restrictions for these constitutive elements. Therefore, their use must comply with the appropriate national laws of each Member State regarding their release. However, Council of Europe Resolution CM/Res(2013)9 on 'metals and alloys used in food contact materials and articles', has recommended the specific release limits (SRL) of 5 mg/kg food for aluminum, whereas no limits for Si and Mg was defined (Europe, 2013). Yet, comparing the extensive application of nanoclays, few researches have addressed the release of nanoclay constituents. The summary of previous studies performed on the migration of nanoclay elements is presented in Table 1 (Avella et al., 2005; Simon et al., 2008; Mauricio et al., 2010; Schmidt et al., 2011; Farhoodi et al., 2014; Maisanaba et al., 2014; Huang J.-Y. A4 - Chieng, Y. Y. A4 - Li, X. A4 -Zhou, W., 2015; Echegoyen et al., 2016). It can be pointed out that most of these investigations have been performed on biosourced and/or biodegradable materials which exhibits a high-water sensitivity. In these cases, the contact with liquid food simulant proved to enhance the migration of elements in higher extent than in polyolefins. The investigation shows very low levels of migration of the three measured mineral elements. As it is evidenced, the migration of Si is higher than the other components for all food simulants which is due to the abundance of Si at the clay surface. It could be assumed that dissolved silica may be generated either by removal of edge-exposed tetrahedral groups (surface hydrolysis) or

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by diffusion of silica through the interlayer space of an expanded Mt. There exists a general agreement that dissolution progresses by breaking of bridge oxygen bonds, Si-O-Al, at the crystal edges (Rozalén et al., 2008). That means that Mt dissolve inwards from the edges, which has been confirmed by AFM observations (Bickmore et al., 2001). External stimuli that would likely affect ENM surface bonds include liquid characteristics (pH, ionic strength, and presence of contaminants that promote bonding), temperature, physical abrasion and vibration. These external factors could dislodge ENMs from the food contact material surface and enable them to release to the simulant. Considering the impact of pH, it is widely known that Al is preferentially released in acidic solutions (Cama et al., 2002). In agreement with this statement, no significant release of aluminum from PNC was observed in the food simulants excepted for acetic acid 3%. However, it should be noted that the demonstrated levels represent the worst case as the polymer was undergone the total immersion which implies that the migration level also includes the release from cut edges. The release of the migrant from the edges, as opposed to the comparatively larger surface area of the faces is assumed to be small for thin films of sufficient diameter, but whether this assumption holds for nanoparticles, which may be more likely to be manually dislodged by the cutting process, is not yet clear. The results of element migration illustrated once again the main influence of food simulant on the migration extent. In addition to the difference in concentration, the relative proportion of each element proved to be variable and dependent on the food simulant. Considering the computation of formula weight of Mt, a unit weight of one clay of 367 g.mol<sup>-1</sup> includes 112 g.mol<sup>-1</sup> of Si, 45 g.mol<sup>-1</sup> of Al and only 8 g.mol<sup>-1</sup> of Mg. The fact that the released quantity did not match this elemental distribution demonstrated that Mt do not migrate under undamaged

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414 form but probably undergoes structural modification during the contact with food simulant and 415 suggest a migration of elements in whole or in part under ionic form. 416 This issue is of high importance regarding the safety point of view because of the possible 417 interactions of released elements. For instance, high concentration of aluminum would be 418 neurotoxic and increase the risk of cognitive impairment and dementia (Gillette-Guyonnet et al., 419 2007). However, several studies (Edwardson et al., 1993; Bellés et al., 1998; Parry et al., 1998) 420 claimed that silicon acts as an antidote to aluminum toxicity by reducing the bioavailability of 421 aluminum. These results could further abate the concerns of aluminum release from the nanoclay 422 PNCs. 423 In regard to iso-octane, although there is no tendency for the clay elements to be released in the 424 food simulant, it should be noted that due to the "aggressive" nature of this simulant towards polyethylene, there could be many other components which are leaving the polymer in contact 425 426 with iso-octane. By leaving the polymer, these products could cause the silicate to migrate to the 427 sample surface which eventually facilitate its release to the simulant. 428 Although ICP-MS provides valuable information on the migration of nanoclay elements, it gives 429 no indication about the migration of nanoparticle per se. Theoretically, nanoclay particles have 430 the potential to release from nanocomposites in contact with food. However, the release process 431 of nanoclay particles may be different due to their size difference and specific chemistry. From a 432 kinetic point of view, the migration of Mt from polyolefin material could be considered as 433 negligible. The slow rate of transfer due to the large size of Mt in comparison with the other 434 spherical nanomaterials exhibiting a radius in the order of magnitude of 1 to 4 nm such as nanoparticle of silver, iron, zinc of titanium (Simon et al., 2008; Bott et al., 2014; Störmer et al., 435 436 2017). To the best of our knowledge, only one investigation evidenced the migration of nanoclays in nanoform released from the surface of the polymer film (Echegoyen et al., 2016). In the study by Schmidt et al. (Schmidt et al., 2011), although the particles ranging from 50-800 nm were detected in 95% ethanol food simulant, none of the characteristic clay minerals were detectable after acid digestion and the analysis by ICP-MS suggested that the detected particles were not of nanoclay origin. In addition, even physical and mechanical surface stresses do not seem to induce migration of inorganic particles such as Laponites, whose structure is close to that of Mt, to be of concern (Bott and Franz, 2019). The results of other studies did not show any nanoclay in the extracts, although it could be argued that this is due to the limitations of the technology rather than the absence of nanoparticles (Chaudhry et al., 2008). One of the techniques that could be applied to detect nanoparticles is SEM-EDX. However, knowing the resolution of the EDX probe (~ 1 µm), it should be noted that this method could identify the agglomeration of nanoclay sheets (in the case of probable migration). As several contamination sources could contribute to the migration results, at the first place, the SEM-EDX of nanoclay powder was acquired and shown in Figure 3 which reveals the majority presence of the constituent elements of the clays but also the occurrence of impurities such as iron or chlorine initially present in the commercial fraction. Based on the EDX results, the atomic ratio of Si/Al and Si/Mg are 2.4±0.1 and 17.1±1.2, respectively. However, for the further particle analysis of the simulants which are filtered through Anodisc membranes, Si/Mg ratio was chosen as the marker because the membrane contains aluminum which could contribute as the background in the EDX results. Some of the SEM-EDX results for the particles detected in filtered simulants are demonstrated in Figure 4.

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Although according to the Figure 4 many particles are found in the simulants, Si/Mg ratio does not comply with corresponding ratio in the nanoclay powder. Therefore, either of the conclusions could be obtained accordingly; 1. All the detected particles are external contaminants and the nanoclay in the form of nanoparticle per se has not migrated through simulants or the size of the released particles are less than the detection limit of SEM-EDX (1  $\mu$ m). 2. Some of the demonstrated particles are representative of the agglomeration of the nanoclays, however, the ratio of elements has been changed through the 10 days of contact due to the dissolution of some elements.

#### 3.4 Additive Migration

During the production of food contact materials, several processing additives are used to improve thermal, optical and mechanical properties of plastic materials. Moreover, during their usage in contact with food, some components may be formed due to the decomposition of the plastic packaging. In the case of nanocomposites, some additional factors would take part in the migration of these additives. In order to understand these interactions, it should be specified that how much of these additives could migrate to the food in a "worst case" scenario, and how could the nanoclay affects the migration of these substances. This issue has already been investigated on high-barrier polymers such as PET (Farhoodi et al., 2017) or PA (de Abreu et al., 2010) to demonstrate the beneficial effects of the barrier effects of nanoclays. This question is worth asking for a low-barrier polymer such as LLDPE for which the incorporation of inorganic nanoparticles can differently modulate the inertness properties.

The migration of organic compounds was measured on a panel of 8 low molecular weight molecules representatives of possible migrants in food-packaging (solvent, wetting agent, plasticizer and photo-initiator in UV-radiation) selected on the basis of the Scientific Opinion edited by EFSA on the criteria to be used for safety evaluation of a mechanical recycling process. The migration levels of the selected substances measured into the recommended food simulants according standard EU conditions (10 days at 40°C) are reported in Figure 5 as the percentage of the migrated additive to FSL divided by the initial quantity of the additive in sample. It is recognized that the release process in the packaging system is controlled by both thermodynamics and kinetics, or partition and diffusion, respectively. Partition coefficient of the migrant between the pure polymer and food (or FSL) at equilibrium status of the migration is affected by the interaction (or affinity) of the migrant with the two phases (polymer and food simulant). For the nanocomposite, the equilibrium status also depends on the additive interactions with compatibilizer (in which the polar fractions alter the nature or chemical properties of the nanocomposite), as well as the nanoclay which could contribute to two different effects: the adsorption sites on the surface of the nanoclay could alter the partitioning of additives physically, whereas the long alkyl chains of the nanoclay modifier could chemically affect the additive partition coefficients. Another aspect of the additive migration from nanocomposite is the kinetics (apparent diffusivity) by introducing a path tortuosity when diffusing substances are forced to migrate around impermeable fillers. The observed results evidenced the major influence of the nature of FSL on the migration process. Regardless of the type of the molecule and sample, the additives were released almost entirely to iso-octane and ethanol 95%. This is in accordance with a previous investigation on the

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diffusivity of the same additives in these two simulants which were already seen to be independent of the type of the molecule (Nasiri et al., 2016). When the polymer is in contact with the fatty food simulants as iso-octane and ethanol 95%, the simulant would be able to penetrate the polymer matrix and interact with the polymer as well as the embedded nanoclay particles and act as a solvent removable penetrant. As a result, the polymer network is swelling which increases its elasticity and eventually accelerates the migration of additives. Moreover, in extreme conditions, penetration of the simulant may lead to the rearrangement of nanoclay and probable movement of nanoclay sheets within the polymer matrix. On the other hand, the interaction of the simulant with nanoclay could lead to the release of the adsorbed additives on the clay surface. As witnessed in Figure 5, in contact with the "non-aggressive" simulants, the release of additives in nanocomposite is either equal or less than the corresponding values in the pure polymer. In these conditions, nanoclay could hinder the additive migration by slowing down diffusion through tortuous path in one hand and the adsorption of additives on the hand. Significance of the either of the effects, depends on the variety of the parameters, such as the polymer-nanoclay interaction (exfoliation, intercalation), size and steric hindrance of the diffusing molecule in addition to the nature of additives, polymer, intercalants and food simulants. As inferred from the results of this research, for a model polymer of poor barrier properties and intrinsic non-favorable nanoclay-LLDPE interactions, and for a selection of worst-case additives, the nanocomposite of LLDPE-compatibilizer-modified nanoclay imparts no risk in terms of the exposure to interactions by-products, migration of intercalants and release of metallic elements.

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#### Conclusion

This study is an attempt through a thorough analysis of the potential risks of a model nanocomposite in food packaging applications by projecting the various safety aspects in a model nanocomposite material, providing a database of the transfer properties of the diverse components of a nanocomposite packaging in common usage conditions and introducing a methodology to optimize the benefit-risk balance of their current usage as food contact materials. As the wide spread application of nanocomposites in food packaging has raised concerns, more studies are directed through the safety evaluation of these materials and most of these researches are limited to either the extent of toxicity as a function of NP type and size or the possibility of NP migration in diverse condition. This investigation presents a broader view over the nanocomposite packaging safety by considering all possible interactions between the polymer, nanoparticle, food, compatibilizer agent, organic modifier and the processing aids or additives and demonstrates that the exhaustive study of the contact suitability of nanocomposite materials is more complex than the evaluation commonly carried out for conventional materials. Assessment of the interaction of nanoclay with some surrogate representatives of the packaging additives or contaminants which was analysed on the basis of the relative migration of these additives revealed that nanoclay tend to hinder the release of these substances in contact with non-aggressive (i.e. aqueous) food simulants while their effect is more contrasted according to the chemical nature of the migrating substance in the case of contact with a fatty simulant. The role of the food simulant nature (or the affinity between polymer and food simulant) remains therefore the dominant factor impacting the mass transfer properties of nanocomposite packaging and, as consequence, the migration extent. In the case of low-barrier polymers such as polyolefins, the migration value measured under standard conditions after 10 days of contact assumes that the system is in an equilibrium state. As consequence, the transfer of additives is

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mainly controlled by thermodynamic rather than kinetic factors. While nanofillers are likely to produce a tortuosity effect and thus limit the diffusion of low molecular weight molecules, their effect on the final measure of specific migration is exerted by a change in the affinity of the migrating molecules for the nanocomposite packaging. The influence of the loading rate and the spatial distribution of nanoclay are then factors that undoubtedly deserve to be investigated in any depth.

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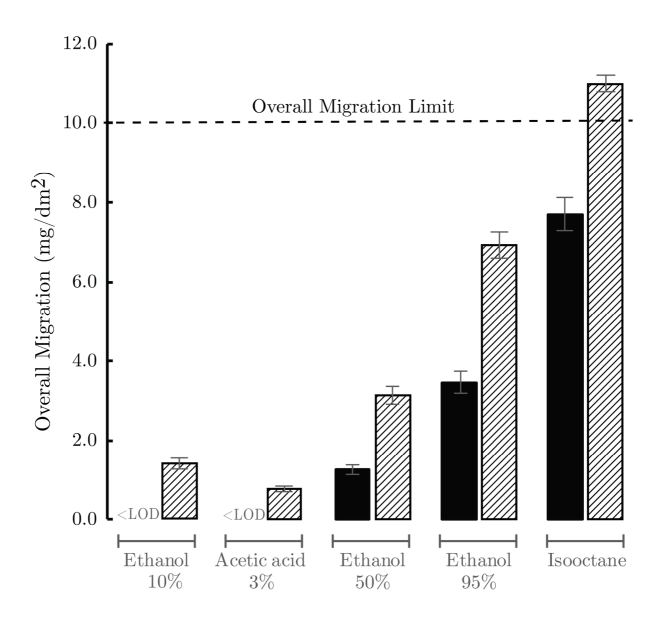
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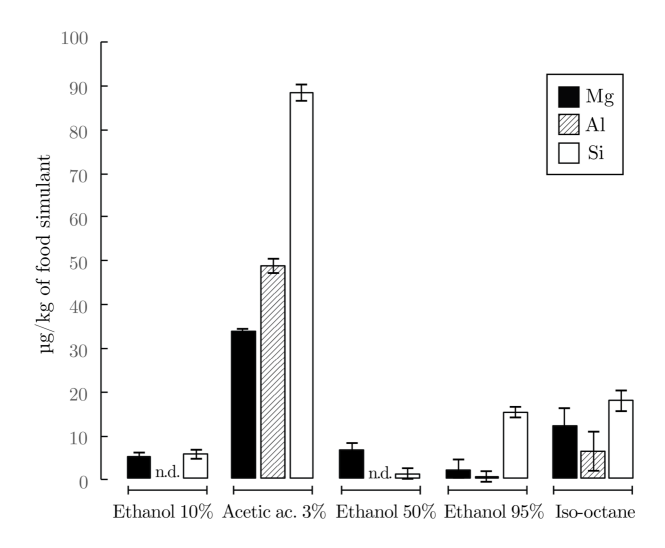
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**Figure 1** Comparison of the overall migration of LLDPE ( ■ ) and PNC ( ② ) to various food simulants (https://doi.org/10.5281/zenodo.3865149)



**Figure 2 :** Migration of Si, Al and Mg from Nanocomposite material to various food simulants (<a href="https://doi.org/10.5281/zenodo.3865149">https://doi.org/10.5281/zenodo.3865149</a>)

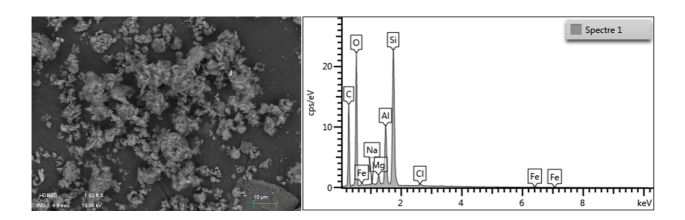
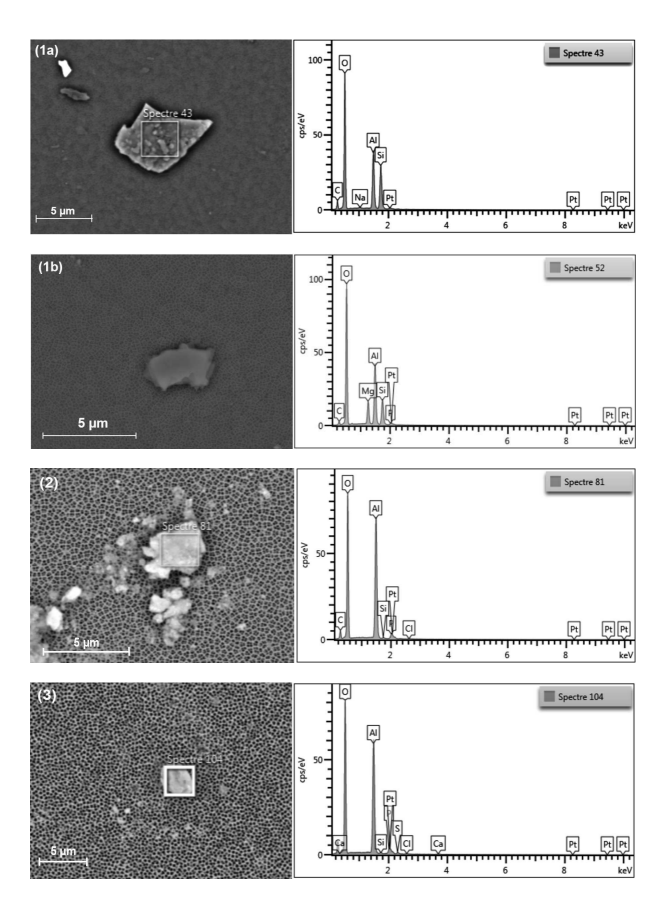
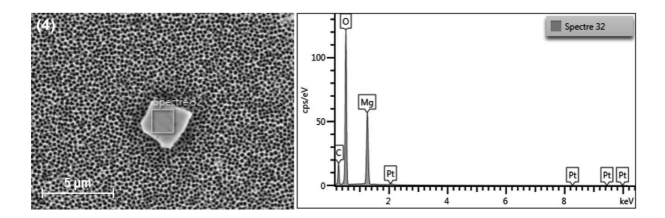


Figure 3 SEM-EDX of nanoclay powder





**Figure 4** SEM-EDX images and corresponding spectra of filtrated particles from iso-octane (1a) and (1b); ethanol 95% (2); ethanol 50% (3) and acetic acid 3% (4).

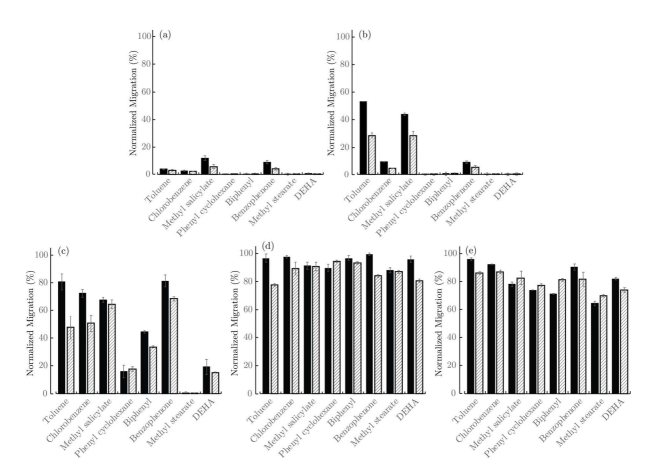


Figure 5 Normalized migration percentage of the selected additives from LLDPE (■) and PNC (□) to Ethanol 10% (a), Acetic Acid 3 w/v% (b), Ethanol 50% (c), Ethanol 95% (d) and Iso-Octane (e) (https://doi.org/10.5281/zenodo.3865149)

**Table 1.** Migration of modifier and compatibilizer from the PNC to the food simulants (mg substance per kg food simulant)

Migrant	Food simulant							
riigi uni	Ethanol 10%	Acetic Acid 3%	Ethanol 50%	Ethanol 95%	Iso-octane			
Maleic Anhydride	< LOD	< LOD	$3.91 \pm 0.97$	8.99 ± 0.99	$0.59 \pm 0.16$			
Quaternary Alkyl Ammonium	$0.01 \pm 0.01$	$0.01 \pm 0.00$	$0.96 \pm 0.13$	$7.6 \pm 0.15$	$3.35 \pm 0.12$			

Table 2. Summary of the researches on nanoclay migration from FCMs according conventional migration testing conditions

Polymer	Food / FSL	Nanoclay type/charge	Migration Condition	Area/Vol (dm²/L)	Migration (mg/kg)	Reference
Starch films Starch / Polyester films	lettuce & spinach	4 w/w %	40°C (10 d)	ND <sup>a</sup>	Si: 16 Mg :260 Si :1.3	Avella et al. 2005
Commercial 3-layer PET bottles	3% HAc	0.2 w/w % <sup>b</sup> Na <sup>+</sup> -MMT	100°C (4 h)	25	None	Chaudhry et al. 2008
PLA films	95% ethanol	5 w/w % Cloisite 30B	40°C (10 d)	ND	Mg: None	Schmidt et al. 2009
Wheat gluten/nanoclay films	water 3% HAc 15% ethanol olive oil	5 w/w % Na <sup>+</sup> -MMT	40°C (10 d)	6	Si: 1.2 (water) Si: 3.5 (3% HAc) Al: 1 (3% HAc) Si: 0.3 (15% ethanol) Si: 0.5 (olive oil)	Mauricio- Iglesias et al. 2010
PET bottles	3% HAc	3 w/w % Cloisite 20A	25°C (90 d) 45°C (90 d)	ND ASTM D- 4754	Al: 0.18, Si: 6 Al: 0.34, Si: 9.5	Farhoodi et al. 2014
PLA composites	distilled water	4 w/w %  MMT/HDTA  4 w/w %  MT/HDTA/ACO	40°C (10 d)	8.2	Mg: 3.6 E-3 Mg:1.51 E-3	Maisanaba et al. 2014
PP/Nanocomposite/PP films	Water 3% HAc 15% EtOH olive oil grapeseed oil coconut oil	4 w/w % MMT	40°C (10 d) °	2	Si:0.12 Si: 1.00 Si: 0.68 Si: 1.50 Si: 1.68 Si: 1.56 Al < LOD	Huang et al. 2015
Aisaika bags (LDPE) Debbie Meyer bags (LDPE)	Ethanol 10% Acetic acid 3%	ND	40°C (10 d) °	16.7	Al: 0.09 Al: 0.04	Echegoyen et al. 2016

<sup>&</sup>lt;sup>a</sup> Not determined (ND)
<sup>b</sup> Averaged over 3 layers

<sup>&</sup>lt;sup>c</sup> Migration data also available for other temperature/time of contact

