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Using FT-IR as a fast method to measure fatty acid soaps in *in vivo* and *in vitro* digests

Kelly Muijlwijk, † Anouk L. Feitsma †* and William Kloek

The objective of the current study was to develop a simple method to measure fatty acid soaps, making use of FT-IR, representative for the soap formation observed in clinical trials. Calcium soaps have a unique coordination which leads to a typical double-splitting of the antisymmetric and symmetric carboxylate peaks. Absorbance values of these carboxylate peaks were used together with the absorbance of the hydrocarbon $-\text{CH}_2$ antisymmetric and symmetric peaks to calculate the calcium soap absorbance. Based on the linear correlation between the calcium soap absorbance and the calcium soap concentration measured with GC-FID, a model was set-up and subsequently successfully validated to quantify calcium soap concentrations in faecal samples from clinical trials with this FT-IR method. With *in vivo* as well as *in vitro* digestion an inverse correlation between the long chain saturated fatty acid part of milk fat containing fat blends used for the infant formulas, and the formation of fatty acid soaps after digestion and defaecation could be observed. There is a clear link between the amount of long chain saturated fatty acids at the *sn*-1/3 position and their release as free fatty acid after lipolysis with the appearance of fatty acid soaps. These insights enable future development of fat blends for infant nutrition to optimize fatty acid soap formation and thereby gut discomfort in infants. These insights can be used to predict the soap formation capacity of a newly designed fat blend and thereby the improvement of infant nutrition products.

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1. Introduction

Human milk is designed by nature and is the best nutrition for healthy growth and development of babies. The fat fraction of human milk is particularly important for a developing infant. It delivers a large part of the required energy as well as essential fatty acids, fat-soluble vitamins and other bioactive components, such as phospholipids and cholesterol.¹ Most of the human milk fat fraction comprises triglycerides with a specific regional distribution of the fatty acids (FA) over the glycerol backbone. Long-chain saturated fatty acids (LCSFAs), *i.e.*, $\geq \text{C12:0}$ esterified at the *sn*-2 position, are easily absorbed during digestion in the form of monoglycerides. In contrast, LCSFAs esterified at the *sn*-1 or *sn*-3 positions (*sn*-1/3), are poorly absorbed after enzymatic hydrolysis because of their low water solubility. Consequently, in the alkaline environment of the small intestinal lumen, free LCSFAs interact readily with cations such as calcium to form insoluble soaps^{2,3} that are associated with hard stools, gut discomfort and decreased absorption of palmitic acid (PA) and minerals by the infant.³⁻⁵

Different fat blends used for infant nutrition, *e.g.*, vegetable oils from different origin and bovine milk fat, contain different levels

of LCSFAs and these fatty acids are also differently distributed on the glycerol backbone. Palmitic acid, the most common LCSFA present in human milk and often also in infant nutrition, is positioned at the *sn*-2 position at 10–20% for commonly used vegetable oil blends and up to 40–45% when vegetable oils and milk fat are combined, while in human milk almost 70% of the palmitic acid is positioned at the *sn*-2 position.^{1,6}

For the measurement of fatty acid soaps in faecal samples, a validated method based on GC-FID has been described by Quinlan *et al.*⁵ Since this is a method that is quite elaborate and specialized, involving multiple extraction steps to isolate the soaped fatty acids, an alternative method that can be performed on small samples in a relatively easy and cost-effective way was explored in the current study.

Fourier transform infrared (FT-IR) spectroscopy is a technique that can be used to measure the chemical composition in small sample amounts, and is for example also used on clinical samples such as body fluids, cells, tissues, nails and faecal samples.⁷ One of the possibilities with FT-IR is a detailed faecal fat analysis,⁸ and it is possible to measure various types of lipids including fatty acid soaps.⁹ Calcium soaps have a typical double-splitting of the antisymmetric carboxylate band at 1575–1580 and 1540–1542 cm^{-1} .^{10,11} This double-splitting is unique for calcium soaps because of the coordination characteristics of calcium when attached to two carboxylate groups.¹¹

FrieslandCampina, Stationsplein 1, 3818 LE Amersfoort, the Netherlands. E-mail: anouk.feitsma@frieslandcampina.com

† Shared first authors.



A clinical trial recently conducted showed that an infant formula containing a blend of vegetable fats and oils and bovine milk fat (50%) reduced the formation of faecal fatty acid soaps and improved stool consistency compared to an infant formula with 100% vegetable fats and oils.¹² Faecal samples obtained in this study were used here to compare the fatty acid soap levels measured with the GC-FID method to levels measured with the FT-IR method.

The objective of the current study was to develop a simple method to measure fatty acid soaps, representative for the soap formation observed in clinical trials. Therefore, we set-up a new method based on FT-IR and correlated these to the soap levels measured with the GC-FID method for faecal soap analysis in the faecal samples obtained from Looijesteijn *et al.*¹³ Next to this, FT-IR measurement of fatty acid soaps formed after *in vitro* digestion was explored. This method would enable the screening of different fat blend compositions for soap formation capacity *via in vitro* digestion and select the most optimal fat blend for clinical testing.

2. Materials and methods

2.1 Measurement of fatty acid soaps in faecal samples

Faecal samples from 3 infant studies were used. For the development of the FT-IR model samples from the study described in Looijesteijn *et al.*¹³ were used. This clinical trial was a single-blinded, crossover, randomized, placebo-controlled study conducted with 19 healthy, full-term, exclusively IF-fed infants.

For the validation of the model, samples from two other studies were utilized. The first set of samples were derived from the clinical trial described in Manios *et al.*¹² This was a double blind, randomized, cross-over trial with 16 healthy, term, formula-fed infants aged 9–14 weeks. The second infant study from which samples were used for validation of the model was a monocentric, randomized cross-over, double blind, referent controlled clinical trial registered as NL7047 in the Dutch trial register. All studies were approved by the local medical ethical committees and parents or caregivers of all participants signed informed consent forms. The studies were conducted in accordance with the guidelines of the Declaration of Helsinki and the International Conference on Harmonisation (ICH) guidelines on Good Clinical Practice (GCP).

Faecal samples were collected during the last three days of each of the intervention periods in faecal tube collector and immediately stored in a home freezer. Transport of the faecal samples was on dry ice. Samples were analysed at Eurofins (Madison, USA) as described below, based on the method of Quinlan *et al.*⁵ Prior to analysis of the stool samples, the samples of each subject were thawed, pooled, homogenized, and lyophilized, and the moisture content was determined gravimetrically. The dried samples were extracted by solvent reflux to obtain the neutral lipids, including non-soaped FFAs. The remaining samples were treated with acetic acid to release the soaped FA which were isolated by a second solvent reflux step. The FFAs were isolated using solid phase extraction. The FFAs were then converted to methyl esters using methanolic hydrochloric acid. The resulting FA methyl esters were analyzed

using a gas chromatograph equipped with a flame ionization detector and quantitated using external standards. Total FA soaps were calculated from the sum of all measured individual FA soaps. Both FFAs and soaped FAs were reported as mg g⁻¹ dry weight stool in the acid form.

Remaining lyophilized faecal samples were stored in a -80 °C freezer and treated with γ -radiation of 40 kGy at Synergy Health (STERIS AST, The Netherlands) to ensure safe handling of the samples before further analysis with FT-IR.

2.1.1 Infant formula and emulsions used for *in vitro* digestion.

The composition of the infant formula that were used for *in vitro* digestion are reported by Manios *et al.*¹² Based on the FA composition the molar percentage of LCSFAs (C12:0-C18:0) positioned at the *sn*-1/3 position on the FAs present was calculated. And with eqn (7) the soap forming capacity was calculated, which is the molar proportion of fatty acids expressed on total fatty acids that are able to form soaps (Table 1). Test formulas were prepared by adding 13.0 g infant formula powder to 90 mL demineralized water and heated to 37 °C in a water bath.

The emulsions used for *in vitro* digestion were prepared to mimic the macronutrient composition of infant formula. Demineralized whey, milk permeate and skim milk were mixed in a ratio of 21 : 57 : 21 and this milk matrix was heated to 50 °C. The fat blends, of which the LCSFA composition is indicated in Table 2, were heated in a water bath at 70 °C until all fat was melted after which the water bath temperature was kept at 60 °C. The fat blends were added to the milk matrix to a final concentration of 3.5 wt% fat. The emulsions were homogenized in 3 rounds with a table top homogenizer at 130 bar.

2.2 *In vitro* digestion

To simulate the digestive system of an infant, a two-phase static *in vitro* digestion model was used, including a gastric and duodenal phase according to the method described by Menard *et al.*¹⁴ The original method was changed as there was no addition of CaCl₂ during the gastric and intestinal phase.

To simulate the gastric phase, 31.5 mL of test formula or home-made emulsion was mixed with simulated gastric fluid (SGF; 18.5 mL) at 525 rpm stirring speed with a magnetic stirrer. SGF was prepared by dissolving rabbit gastric extract (RGE; CNRS, Marseille, France) at 0.6 mg mL⁻¹ (with a lipase conc of 46 μ g mL⁻¹) in a 13 mM KCl and 94 mM NaCl solution. RGE contains rabbit gastric lipase and was chosen because of similar properties to human gastric lipase, like regional- and stereo-chemical preference.¹⁵ RGE was added to the gastric juice to reach a final

Table 1 Infant formula that varied in milk fat content and their molar percentage of C12:0-C18:0 LCSFA positioned at *sn*-1/3 and the calculated soap forming capacity (SFC) of these infant formulas

Milk fat (%)	LCSFA <i>sn</i> -1/3 (mol%)	SFC
50	32.0	21.3
20	48.3	32.2
0	52.0	34.7



Table 2 Emulsions with different fat blends used for *in vitro* digestion with their molar percentage of C12:0–C18:0 LCSFA positioned at *sn*-1/3 and the calculated soap forming capacity (SFC)

Sample	LCSFA <i>sn</i> -1/3 (mol%)	SFC
1	30.0	20.0
2	32.0	21.3
3	37.8	25.2
4	52.0	34.7

gastric lipase concentration of 17 $\mu\text{g mL}^{-1}$ in the digestion mixture, representative for the gastric lipase concentration in gastric contents at half gastric emptying in infants.¹⁶ During addition of gastric juice, pH is adjusted to 5.5 with 1 M HCl. During the gastric phase the pH was kept constant at pH 5.5 by addition of 0.1 M HCl and/or 0.1 M NaOH. Products with gastric juice were incubated for 60 min at 37 °C. Then the pH was adjusted to 6.5 with 0.1 M NaOH and 27.6 mL of intestinal juice containing bovine bile salts (3.1 mM; Sigma-Aldrich, St Louis, USA), 10 mM KCl and 41 mM NaCl was added and incubated for 15 minutes. After that porcine pancreatic extract (90 U mL^{-1} ; PPE or pancreatin; 8xUSP, Sigma-Aldrich, St Louis, USA) was added and incubated for 1.5 hours. During the intestinal phase the pH was kept at pH 6.5 using 0.1 M NaOH. The enzymatic reaction of all samples was inhibited by directly cooling the samples on ice. The digests were centrifuged at 13 500g for 30 minutes at 7 °C and the pellets were freeze-dried and stored in a –20 °C freezer until further measurement. The typical weights of the pellets were 0.4–0.9 g after centrifugation and 0.04–0.16 g after freeze-drying. Digestion of the test formulas was performed in single fold and digestion of emulsions in triplicate.

2.3 FT-IR

Samples obtained after *in vivo* (Section 2.1) and *in vitro* (Section 2.2) digestion were measured in duplicate with FT-IR. As a reference for calcium soaps calcium dipalmitate (Sigma-Aldrich) was measured. Attenuated total reflectance (ATR) spectra were obtained using the build-in diamond ATR of the Nicolet iS50 FT-IR Spectrometer (ThermoFisher Scientific) equipped with a DTGS detector and a KBr beamsplitter. Before each measurement a background scan was performed with air. Freeze-dried samples were grinded to ensure homogeneity. A pressure of 27 kg was applied with the build-in pressure device on a sample of 10–30 mg that was placed on the all-reflective diamond ATR. Spectra were measured at wavenumbers ranging from 4000 to 400 cm^{-1} with a resolution of 4 cm^{-1} and an accumulation of 64 scans. Collected spectra were analyzed using Omnic 9 (ThermoFisher Scientific). A linear baseline correction was performed with baseline points at 4000, 3800, 2400, 1900, 825 and 400 cm^{-1} using the Auto Y function. Peak height and position at peak apex was measured with the Find Peaks function.

3 Results and discussion

3.1 Calcium soaps in *in vivo* digests

3.1.1 FT-IR peak assignment of calcium soaps in *in vivo* digests. The FT-IR spectra of calcium dipalmitate and a representative faecal sample are depicted in Fig. 1. Assignment of relevant peaks is given in Table 3. Calcium soaps have a typical peak pattern due to the carboxylate group attached to the calcium ion. Fig. 1 shows that the antisymmetric (1575 and 1538 cm^{-1}) and symmetric (1434 and 1420 cm^{-1}) COO^- stretching peaks are both present as a dimer. These dimers indicate that the calcium ion coordination structure has two different denticities, most likely in the bidentate and unidentate

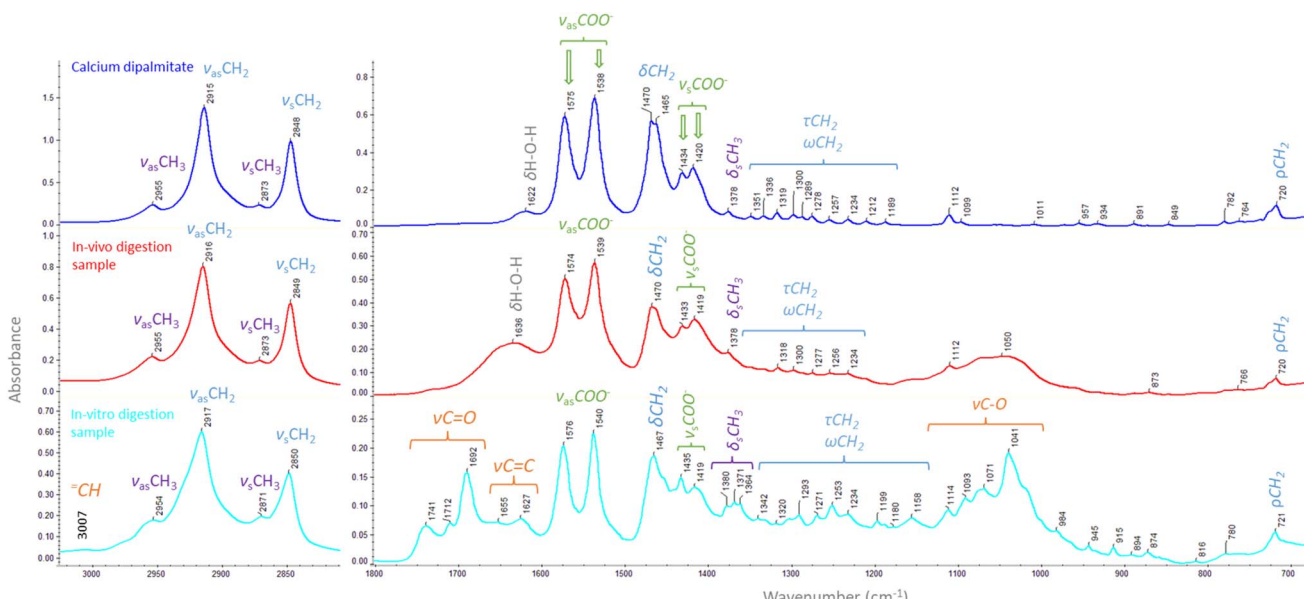


Fig. 1 FT-IR spectra of calcium dipalmitate (blue), a representative *in vivo* sample (red), and a representative *in vitro* sample (cyan) with assigned peaks in Table 3.



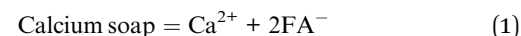
Table 3 Assignment functional groups in FT-IR spectra of infant formula after *in vivo* and *in vitro* digestion

Wavenumbers (cm ⁻¹)			
<i>In vivo</i>	<i>In vitro</i>	Assignment	Notes
3600–3000		νOH	Stretching of O–H
	3008–3007	$=\text{CH}$	C–H symmetric stretching in a cis double bond
2956–2954	2955–2954	$\nu_{\text{as}}\text{CH}_3$	C–H antisymmetric stretching of –CH ₃
2917–2915	2918–2917	$\nu_{\text{as}}\text{CH}_2$	C–H antisymmetric stretching of –CH ₂
2873–2872	2872–2871	$\nu_s\text{CH}_3$	C–H symmetric stretching of –CH ₃
2849–2848	2851–2849	$\nu_s\text{CH}_2$	C–H symmetric stretching of –CH ₂
	1741–1735	$\nu\text{C=O}$	TAG ester
	1712		Saturated FFA
	1692–1691		Unsaturated FFA
	1655, 1627	$\nu\text{C=C}$	
1637–1630		$\delta\text{H–O–H}$	Scissoring of H–O–H
1575–1574	1583–1576	$\nu_{\text{as}}\text{COO}^-$	Antisymmetric stretching of –COO [–]
1540–1538	1542–1540		
1472–1466	1468–1465	δCH_2	Scissoring of –CH ₂
1435–1431	1436–1435	$\nu_s\text{COO}^-$	Symmetric stretching of –COO [–]
1420–1417	1419–1411		
1380–1378	1380–1379	$\delta_s\text{CH}_3$	Symmetric scissoring of –CH ₃
	1371–1370		
	1365–1364		
1320–1230	1350–1150	τCH_2	Twisting of –CH ₂
		ωCH_2	Wagging of –CH ₂
	1120–1000	$\nu\text{C–O}$	Stretching of esters
720–719	721–719	ρCH_2	Rocking of –CH ₂

coordination.¹⁰ The location of the $\nu_{\text{as}}\text{COO}^-$ peak at 1575 cm⁻¹, as well as the splitting value of ~40 cm⁻¹ between the antisymmetric peaks, confirm the presence of calcium soaps.¹⁰

The hydrocarbon tail of calcium soaps gives rise to –CH₂ and –CH₃ peaks in the region 2955–2849 cm⁻¹ as well as a peaks at 1470, 1378–1190, and 720 cm⁻¹.¹⁰ The absence of C=O stretching carbonyl peaks typical for fatty acids¹⁰ at 1703 cm⁻¹ indicate that the hydrocarbons present in the sample can all be attributed to those present in calcium soaps.

3.1.2 Model definition. Calcium soaps are composed of a calcium ion linked *via* carboxylate groups with two fatty acid tails: calcium soap = FA₁–Ca–FA₂ = R₁–COO₁–Ca–COO₂–R₂ (eqn (1)). The solubility product (K_{sp}) of calcium soaps is presented in eqn (2). Based on the solubility product the calcium soap response in an FT-IR measurement is defined in (3) and consists of the calcium part, represented by the attached carboxylate groups, multiplied with the FT-IR response for the fatty acid tails. The carboxylate FT-IR response for both carboxylate groups together is the sum of the two carboxylate symmetric stretching vibration peaks at 1434 and 1420 cm⁻¹, and the two carboxylate antisymmetric stretching vibrations at 1575 and 1538 cm⁻¹. The FT-IR response of the two fatty acid tails are combined as the sum of the CH₂ symmetric and antisymmetric stretching vibrations at 2848 and 2915 cm⁻¹. Combining these specific peaks for the carboxylate and fatty acid tails resulted in a definition for the calcium soap absorption response ($A_{\text{Ca-soap}}$) (eqn (4)). In this equation the absorption of the individual peaks is the peak height as measured after baseline correction.



$$K_{\text{sp}} = [\text{Ca}^{2+}] \times [\text{FA}^-]^2 \quad (2)$$

$$A_{(\text{Ca-soap})} = (\text{COO}_1^- + \text{COO}_2^-) \times R^2 \quad (3)$$

$$A_{(\text{Ca-soap})} = (\nu_s\text{COO}_1^- + \nu_{\text{as}}\text{COO}_1^- + \nu_s\text{COO}_2^- + \nu_{\text{as}}\text{COO}_2^-) (\nu_s\text{CH}_2 + \nu_{\text{as}}\text{CH}_2)^2 \quad (4)$$

We used 31 samples from a previous study¹³ to set-up a model to calculate the calcium soap content in *in vivo* samples based on FT-IR measurements. Samples were measured with FT-IR and resulting absorption peak heights were used to calculate $A_{\text{Ca-soap}}$ with eqn (4). The calcium soap concentration in these samples was measured in the study of Looijesteijn *et al.*¹³ with GC-FID and were plotted against the $A_{\text{Ca-soap}}$ as measured in the present study with FT-IR (Fig. 2). Linear regression showed that there was a linear relation through the origin between calcium soap absorption and the calcium soap concentration ($C_{\text{Ca-soap}}$, in mg g⁻¹ dry faeces) (eqn (5)). The fitting parameter c was 93.421 mg g⁻¹ with a 95% confidence interval of 89.843 to 97.000 and a *p*-value of <0.0001. This linear relation can be further used to calculate the calcium soap concentration in *in vivo* samples *via* FT-IR measurements.

$$C_{\text{Ca-soap}} = c \times A_{\text{Ca-soap}} \quad (5)$$

3.1.3 Model validation. To validate the calcium soap model, two different validation sets were used. From Manios *et al.*¹² 32 faecal samples from their cross-over study 1 were



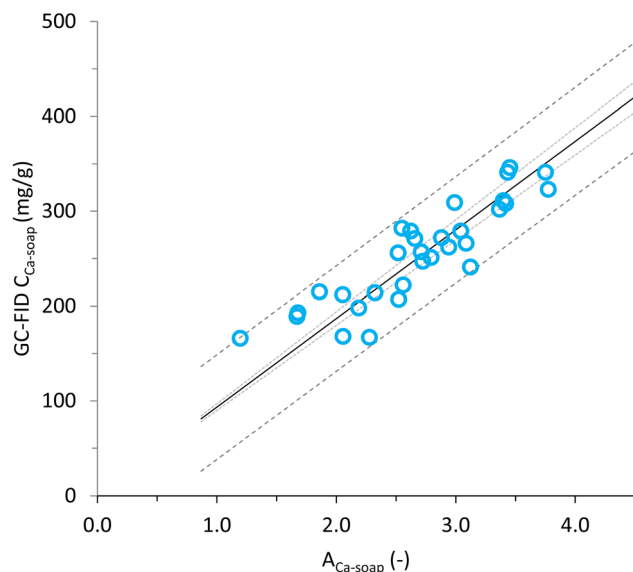


Fig. 2 Fit between absorbance of calcium soaps ($A_{\text{Ca-soap}}$) as measured with FT-IR and calcium soap concentration ($C_{\text{Ca-soap}}$) as measured with GC-FID, on dry faeces samples from the study described in Looijesteijn *et al.*¹³ With the model (—), 95% confidence interval on mean of prediction (···) and 95% confidence interval on single prediction (—).

used. Another set of 9 faecal samples from an infant feeding study (registered as NL7047 in the Dutch trial register) was used. Absorbance heights of the carboxylate and fatty acid peaks of these faecal samples were measured with the FT-IR and used to calculate $A_{\text{Ca-soap}}$ with eqn (4). With the earlier defined model in eqn (5) the calcium soap concentration ($C_{\text{Ca-soap}}$) was calculated. To validate the model, the $C_{\text{Ca-soap}}$ measured with FT-IR was plotted against $C_{\text{Ca-soap}}$ as measured with GC-FID (Fig. 3A). Prediction of the calcium soap concentration based on FT-IR measurements was for both independent validation sets within the 95% confidence interval. A Bland-Altman¹⁷ analysis

was performed to evaluate the agreement between both methods to measure calcium soap concentration (Fig. 3B). In this plot the difference in calcium soaps as measured between the GC-FID and FT-IR is plotted against the average of the measurement. The mean difference is -6.7 mg g^{-1} dry feces with a 95% confidence interval of -14.0 to 0.6 mg g^{-1} dry feces. In the plot are also the limits of agreement, which are defined as the range within 95% of the differences between the measurements would occur. All measurement points fall within the limits of agreement from which we can conclude that the FT-IR method is in agreement with the GC-FID method. Based on the plot we can also conclude that there is no relationship between the differences and the range of the true value. This indicates that the FT-IR method is a reliable method to quantify fatty acid soaps and is only marginally influenced by the parameters that were possibly different between the clinical studies, *e.g.*, collection and transportation of samples, faecal composition and product composition differences in the test products used in both intervention studies.

3.2 Calcium soaps in *in vitro* digests

3.2.1 FT-IR measurement on *in vitro* digests. *In vitro* digestion samples are generally too small to measure calcium soap concentration with the GC-FID method.⁵ Sample size is sufficient for an FT-IR measurement and the method as set-up previously for *in vivo* samples can thus potentially be used to obtain an indication of calcium soap formation during digestion. In this paragraph the measurement of calcium soaps in *in vitro* samples using FT-IR is discussed.

3.2.1.1 Qualitative interpretation of the FT-IR spectra. A comparison of spectra after *in vivo* and after *in vitro* digestion is presented in Fig. 1. Assignment of peaks relevant for calcium soap quantification is largely consistent with those assigned for the *in vivo* samples, albeit that there are some slight shifts in wavenumber at which the peaks appear (Table 3). Next to shifts also some additional peaks appear in the *in vitro* spectra. A C-H

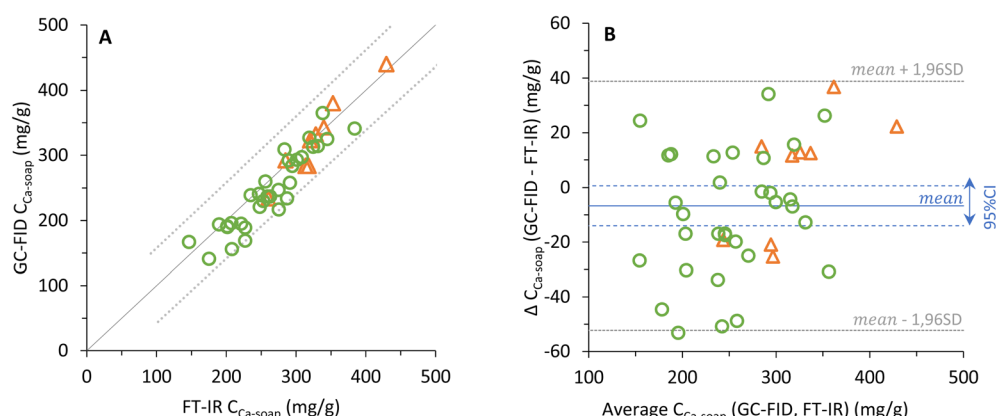


Fig. 3 (A) Calcium soap concentration in dry faecal samples from Manios *et al.*¹² (○) and from an infant feeding study (registered as NL7047 in the Dutch trial register) (Δ) measured with GC-FID plotted against the calcium soap concentration based on FT-IR measurements, with the parity line (—) and 95% confidence interval (···). (B) Bland-Altman¹⁷ plot of the same data in which the difference against mean is plotted for calcium soap with the mean difference (\bar{d} , —) and its 95% confidence interval (—) and the limits of agreement ($\bar{d} \pm 1.96 \cdot \text{standard deviation of the differences}$, ···).



symmetric stretching vibration of the cis double bond appears at 3012 cm^{-1} .¹⁸ The peaks present at 1741 , 1712 and 1692 cm^{-1} can be assigned to vibration of carboxylic acids. The peak at 1741 cm^{-1} represents the ester functional group in glycerides. The peaks at 1712 and 1692 cm^{-1} represents the C=O group in free fatty acids (FFA), of which the first represents saturated FFAs and the latter unsaturated FFAs.¹⁹ Unsaturated FFAs also give rise to the C=C peak at 1655 cm^{-1} .²⁰ Symmetrical scissoring of the methyl groups has changed from one peak at 1380 cm^{-1} to a triplet at 1380 , 1371 , and 1364 cm^{-1} . Peaks present between 1350 and 1150 cm^{-1} are most likely due to twisting and wagging vibrations of the methylene group.^{21,22} C-O stretching vibrations of esters is represented by the peaks located at 1157 , 1113 , 1070 and 1041 cm^{-1} .^{21,23} Between 1000 and 780 cm^{-1} multiple small peaks appear which can most likely be attributed to methylene rocking and twisting.²²

The extra functional groups that are present in the *in vitro* digest compared to the *in vivo* digest are all lipid related and indicate that there are glycerides present as well as unsaturated and saturated FFAs. During *in vivo* digestion the monoglycerides and short chain FFAs formed during lipolysis are absorbed in the small intestine, as well as minerals such as calcium. The long chain FFAs are either absorbed or precipitated in a calcium soap. While during *in vitro* digestion monoglycerides, short chain FFAs and minerals remain in the sample. At the end of *in vitro* digestion there is a centrifugal separation during which a large part of the lipids are removed with the cream layer and the calcium soaps precipitate in the pellet that is collected. However, this separation is not ideal and thus also glycerides and FFAs end up in the pellet as well as insoluble minerals.

3.2.1.2 Quantitative interpretation of the FT-IR spectra. For each sample the calcium-soap absorption ($A_{\text{Ca-soap}}$) was calculated with eqn (4) and subsequently multiplied with the mass fraction of dry pellet ($m_{\text{dry pellet}}$) to wet pellet ($m_{\text{wet pellet}}$) with eqn (6) to calculate the calcium soap formation index. The mass fraction is required to correct for the amount of pellet that is formed during centrifugation.

$$\text{Calcium soap formation index} = \frac{m_{\text{dry pellet}}}{m_{\text{wet pellet}}} \times A_{\text{Ca-soap}} \quad (6)$$

To test the applicability of FT-IR to measure calcium soap formation during *in vitro* digestion, three model infant formulas containing 0 , 20 or 50% milk fat, were digested in an *in vitro* digestion model. The infant formulas with 0 and 50% milk fat in the fat blend were the same as used in the *in vivo* study¹³ that was used for the model definition.

The concentration of calcium soaps in the *in vivo* faecal samples measured with the FT-IR method decreased for the fat blends with increasing milk fat contribution of the long chain saturated fatty acid fraction in the infant formula (Fig. 4), as was also measured with the validated GC-FID method.¹² Also after *in vitro* digestion an inverse correlation between the fat blends including milk fat contributing in the long chain saturated fatty acid part of the fat blend used for the infant formulas, and the calcium soap formation index was observed (Fig. 5). Note here that the calcium soap formation index obtained with FT-IR

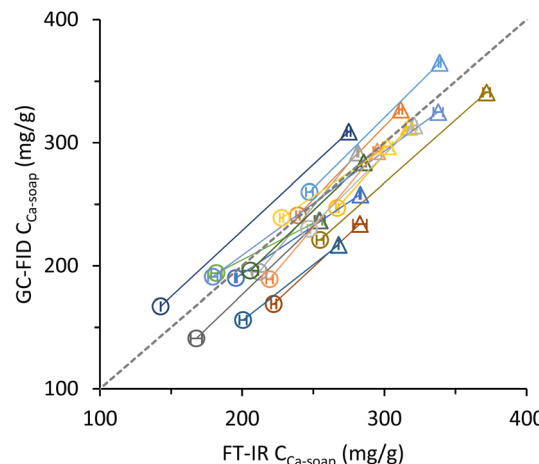


Fig. 4 Fatty acid soap concentration measured with GC-FID as a function of the fatty acid soaps measured with FT-IR in *in vivo* digestion samples from infants fed with infant formula containing 50% (●) or 0% (▲) milk fat in the fat blend of the infant formula.¹² Each line represents the faecal samples of one infant. Error bars represent the standard deviation of duplicate FT-IR measurement.

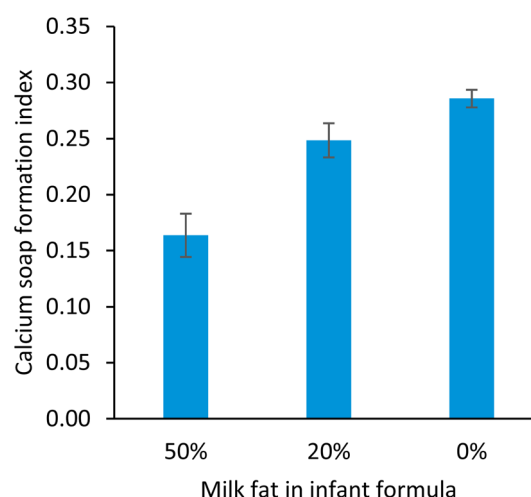


Fig. 5 Calcium soap formation index as measured with FT-IR after *in vitro* digestion of infant formula with varying milk fat content in the fat blend. Error bars represent the standard deviation of duplicate FT-IR measurement.

measurements on *in vitro* digests yields a relative index rather than an exact value of the formed calcium soaps. These results show that the FT-IR method presented in this manuscript can be used to predict relative calcium soap formation of an infant formula during *in vitro* digestion.

3.2.2 Predicting soap formation of fat blends. Because only fatty acids present at the *sn-1/3* position can form calcium soaps, a theoretical soap formation capacity (SFC) can be calculated.²⁴ Kloek *et al.*²⁴ defined the SFC as the mole % of long chain saturated fatty acids present at the *sn-1/3* position on the total mole % of fatty acids present in the fat blend, in which the index i stands for the individual LCSFA (eqn (7)).



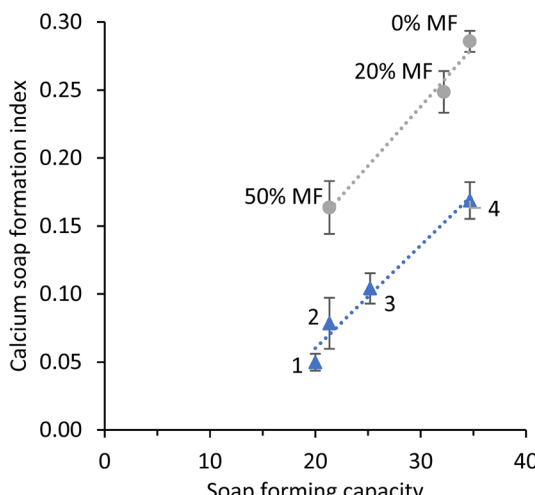


Fig. 6 Calcium soap formation index as measured with FT-IR after *in vitro* digestion of infant formula with varying milk fat content in the fat blend (●) and emulsions containing different fat blends (▲) as a function of the soap forming capacity with a linear trendline fitted through the data. Error bars represent a different standard deviation for each sub-set: For the infant formula (●) duplicate FT-IR measurement was performed on single fold *in vitro* digestion, and for the emulsions (▲) single FT-IR measurement was performed on *in vitro* digestion conducted in triplicate.

$$SFC = \frac{2}{3} \sum_i FA_{sn13,i} \quad (7)$$

Calculation of the SFC can be used to optimise the design of fat blends towards human milk fat, which has a typical SFC of ~ 20 . Kloek *et al.*²⁴ showed that the soap forming capacity had a positive correlation with total long chain saturated fatty acids measured in stools, which originates from the same *in vivo* study as was used in this study.

To investigate whether there is a relationship between the measured calcium soaps in *in vitro* digests and the theoretically calculated soap-forming capacity, the data from the calcium soap formation index presented in Fig. 5 were plotted against the soap formation capacity (Fig. 6). Next to that, a new set of *in vitro* digests was measured with FT-IR and the resulting calcium soap formation index was also plotted in Fig. 6. In both sets the calcium soap formation index had a linear correlation ($R^2 = 0.98$ and 0.97) with the soap forming capacity. However, calcium soap formation index was different at equal soap forming capacity. The shift in calcium soap formation index is due to the effect of the sample matrix on the FT-IR measurement. For example, variation in proteins and minerals affect the amount of pellet formed and thus the calcium soap formation index. It is therefore recommended to compare results with a reference of the same matrix. In this case the set with varying milk fat content can be compared within the set, but not with the set of measurements that consisted of emulsions containing different fat blends. Based on these results we can conclude that for a given matrix the expected calcium soap formation can be steered well by optimizing the fat blend composition with

respect to its regional fatty acid distribution and taking into account specific regulatory requirements.

4. Conclusion

The objective of the current study was to develop a simple method to measure fatty acid soaps, representative for the soap formation observed in clinical trials. Therefore, the newly identified method based on FT-IR was compared to the soap levels measured with the validated method, consisting of an extraction followed by GC-FID detection, for faecal soap analysis in samples of clinical trials performed with infants consuming infant formulas with different fat blends. There was a linear correlation between the calcium soap absorbance, which was calculated with absorbance values of carboxylate and hydrocarbon peaks measured with the FT-IR, and the calcium soap concentration as measured with GC-FID in *in vivo* digestion samples. Using this correlation, a model was set-up with which calcium soap concentration in *in vivo* samples can be measured with a simple FT-IR measurement. This model was validated with *in vivo* samples from another clinical trial performed in infants. The good fit of the model indicates that the FT-IR method is a reliable method to quantify fatty acid soaps and is only marginally influenced by parameters that were possibly different between the clinical studies.

In vitro digestion samples are generally too small for calcium soap quantification *via* GC-FID. We have shown here that calcium soaps can be measured with FT-IR using the same method as for *in vivo* digestion samples. However, for *in vitro* samples, the FT-IR method provides a relative index rather than an exact value of the formed calcium soaps. Most likely this is the case because there is no absorption of released components during *in vitro* digestion and these components may interfere with the calcium soap peaks needed for quantification. Hence, the *in vitro* method is highly matrix dependent, and results can therefore only be compared within a set of samples with limited variation in composition.

With *in vivo* as well as *in vitro* digestion an inverse association between the fat blends used for infant nutrition products containing different vegetable fat blends combined with different amounts of milk fat proportion and the formation of fatty acid soaps after digestion and defaecation could be observed. There is a clear link between the amount of long chain saturated fatty acids at the *sn*-1,3 position and therefore release as free fatty acid after lipolysis with the appearance of fatty acid soaps, providing extra substantiation for the future development of fat blends for infant nutrition to minimize fatty acid soap formation and thereby gut discomfort in infants. These insights can be used to predict the soap formation capacity of a newly designed fat blend and thereby the improvement of infant nutrition products.

Author contributions

Kelly Muijlwijk: formal analysis, investigation, methodology, validation, visualization, writing – original draft, writing – review and editing. Anouk Feitsma: conceptualization, funding acquisition, investigation, project administration, resources,



supervision, visualization, writing – original draft, writing – review and editing. William Kloek: conceptualization, methodology, validation, writing – review and editing.

Conflicts of interest

Authors are all employed by FrieslandCampina. There are no further conflicts of interest to declare.

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