Technical note: Near infrared reflectance spectroscopy to predict fecal indigestible neutral detergent fiber for dairy cows

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ABSTRACT

In vitro and in situ procedures performed to estimate indigestible neutral detergent fiber (iNDF) in forage or fecal samples are time consuming, costly, and limited by intrinsic factors. In contrast, near infrared reflectance spectroscopy (NIRS) has become widely recognized as a valuable tool for accurately determining chemical composition and digestibility parameters of forages. The aim of this study was to build NIRS calibrations and equations for fecal iNDF. In total, 1,281 fecal samples were collected to build a calibration data set, but only 301 were used to develop equations. Once dried, samples were ground and chemically analyzed for crude protein, ash, amylase and sodium sulfite–treated NDF corrected for ash residue (aNDFom), acid detergent fiber, acid detergent lignin, and in vitro digestion at 240 h to estimate iNDF (uNDF240). Each fecal sample was scanned using a NIRSystem 6500 instrument (Perstorp Analytical Inc., Silver Spring, MD). Spectra selection was performed, resulting in 301 sample spectra used to develop regression equations with good accuracy and low standard error of prediction. The standard error of calibration (SEC), cross validation (SECV), and coefficients of determination for calibration (R²) and for cross validation (1 − VR, where VR = variance ratio) were used to evaluate calibration and validation results. Moreover, the ratio performance deviation (RPD) and ratio of the range of the original data to SECV (range/SECV; range error ratio, RER) were also used to evaluate calibration and equation performance. Calibration data obtained on fiber fractions aNDFom (R² = 0.92, 1 − VR = 0.87, SEC = 1.48, SECV = 1.89, RPD = 2.80, and RER = 20.19), uNDF240 (R² = 0.92, 1 − VR = 0.86, SEC = 1.65, SECV = 2.24, RPD = 2.57, and RER = 14.30), and in vitro rumen aNDFom digestibility at 240 h (R² = 0.90, 1 − VR = 0.85, SEC = 2.68, SECV = 3.43, RPD = 2.53, and RER = 14.0) indicated the predictive equations had good predictive value. Key words: undigested NDF, fecal composition, near infrared reflectance spectroscopy

Technical Note

Ruminal digestion of NDF is linked to the part of NDF that is indigestible and the rate at which the potentially digestible NDF is fermented. Indigestible NDF (iNDF; determined by in situ or in vitro procedures) is reported to be a reliable digestibility marker in controlled animal experiments (Lippke et al., 1986; Huhtanen et al., 1994, 2006a; Lund et al., 2007). The standard nomenclature throughout the literature refers to “indigestible NDF” (Mertens, 1993; Huhtanen et al., 2006b). To improve the accuracy of the standard terminology used to describe fiber fermentation dynamics, Mertens (2013) coined the term “undigested NDF” (uNDF) as the laboratory measure of indigestible NDF at a specified fermentation time. The uNDF fraction could be estimated via long-term (240 h) in vitro fermentations (uNDF240; Palmonari et al., 2016) or by incubating the samples in bags placed in the rumen for 288 h. However, this procedure relies on the availability of cannulated cows and is limited by the intrinsic characteristics of the bags used (Huhtanen et al., 2006a; Krizsan et al., 2012). The uNDF240 fraction represents the fiber fraction that affects physical effectiveness, digestion, passage rates, and gut fill of forages (Van Amburgh et al., 2015). Moreover, by measuring fecal iNDF output, we could acquire additional information on the proportion of potentially digestible NDF really digested in vivo. Furthermore, fecal uNDF is useful as an intrinsic digestibility marker compared with total fecal collection (Lee and Hristov, 2013) for a total-tract nutrient digestibility estimation for dairy cows (Ferraretto et al., 2015). According to Palmonari et al. (2016), uNDF240 should be included in routine forage and feed analysis because it offers a more predictable digestibility and uniformity than NDF alone. As reported above, wet
chemical analyses have been used to characterize forages and predict their nutritive value. However, these are time consuming (especially regarding uNDF240 determination) and costly, and may involve the use of hazardous chemicals. Near infrared reflectance spectroscopy (NIRS) has become widely recognized as a valuable tool in accurate determination of the chemical composition of a wide range of forages (Shenk and Westerhaus, 1985; Redshaw et al., 1986) and in the prediction of digestibility parameters (Norris et al., 1976; Barber et al., 1990; Park et al., 1997). A close relationship between NIRS and iNDF of grass silage was reported first by Nousiainen et al. (2004), and robust NIRS calibrations are already available, using C3 and C4 forages, for the prediction of iNDF (Harper et al., 2014). Although several studies have been conducted, no experiments have shown the correlation between NIRS spectra and fecal uNDF of dairy cows, especially using long in vitro incubations in individual batch fermentation systems. The aim of this work was to construct NIRS calibrations and equations for fecal uNDF240, using an in vitro approach, to aid nutritionists to adapt the cow response to forage quality, improving the efficiency of use of forage fiber in diet for dairy cows.

In total, 1,281 fecal samples were collected but only 301 were used to build the calibration. The samples were collected by rectal grab from cows involved in 4 feeding trials. In the first trial (157 original samples, 39 selected for calibration), heifers were fed with dried, ground, and pelleted grass hay of different quality mixed with TMR (Bonfante et al., 2016). In the second trial (548 original samples, 113 selected for calibration), lactating cows were fed a diet based on alfalfa that differed in harvest times and digestibility (Fustini et al., 2014). In the third trial (257 original samples, 25 selected for calibration), lactating cows were fed a diet based on grass hay and alfalfa, mixed with TMR (Bonfante et al., 2016). In the fourth trial (548 original samples, 113 selected for calibration), heifers were fed with dried, ground, and pelleted grass hay of different quality mixed with TMR (Bonfante et al., 2016). Rumen fluid was collected from 2 lactating cows fed a hay-based diet (milk production = 33.2 ± 1.7 kg/d; 251 ± 2 DIM) containing grass hay and alfalfa, corn barley and sorghum as energy source, and soybean meal as protein source (Palmonari et al., 2016). Each sample was analyzed in triplicate. Rumen fluid was collected from 2 lactating cows fed a hay-based diet (milk production = 33.2 ± 1.7 kg/d; 251 ± 2 DIM) containing grass hay and alfalfa, corn barley and sorghum as energy source, and soybean meal as protein source (Palmonari et al., 2016). Rumen fluid was mixed and placed in a CO2-gassed insulated bottle. Rumen contents were filtered through 4 layers of cheesecloth and kept under constant O2-free CO2 flow before inoculating each flask. After 240 h of incubation, sample residues were analyzed to determine the aNDF0m content, following the procedure described by Mertens (2002), which represents the uNDF240 of the fecal samples determined in these conditions. The duration of incubation was based on previous experiences that showed 240 h to be the maximum length for in vitro fiber digestion under anaerobic conditions (Fox et al., 2004; Raffrenato and Van Amburgh, 2010; Cotanch et al., 2014). Indigestible aNDF0m, after a 240-h incubation, is determined after 4-h combustion at 550°C in a muffle furnace, and starch (AOAC, 1990). As a good predictor of uNDF, in vitro rumen aNDF0m digestibility at 240 h (IVNDFD240) was determined using the Tilley and Terry modified technique (Tilley and Terry, 1963; Robertson and Van Soest, 1981), according to the procedure described by Palmonari et al. (2017). Briefly, basic procedures consisted of incubating 0.5 g of dried and ground fecal sample in a 150-mL Erlenmeyer flask under CO2 back-pressure at 39°C in a water bath for 240 h; the flask contained rumen fluid, buffer medium, and macro- and micromineral solution (Goering and Van Soest, 1970). Each sample was analyzed in triplicate. Rumen fluid was collected from 2 lactating cows fed a hay-based diet (milk production = 33.2 ± 1.7 kg/d; 251 ± 2 DIM) containing grass hay and alfalfa, corn barley and sorghum as energy source, and soybean meal as protein source (Palmonari et al., 2016). Rumen fluid was mixed and placed in a CO2-gassed insulated bottle. Rumen contents were filtered through 4 layers of cheesecloth and kept under constant O2-free CO2 flow before inoculating each flask. After 240 h of incubation, sample residues were analyzed to determine the aNDF0m content, following the procedure described by Mertens (2002), which represents the uNDF240 of the fecal samples determined in these conditions. The duration of incubation was based on previous experiences that showed 240 h to be the maximum length for in vitro fiber digestion under anaerobic conditions (Fox et al., 2004; Raffrenato and Van Amburgh, 2010; Cotanch et al., 2014). Indigestible aNDF0m, after a 240-h incubation, is determined after an incineration step at 550°C for 3 h and is expressed on an OM basis as a percentage of DM. The IVNDFD240 (percentage of aNDF0m) was also determined for each fecal sample, from the same assay, using standard calculations (Goering and Van Soest, 1970).

For spectra acquisition, each fecal sample was packed into cylindrical sample holders equipped with a quartz window and scanned (wavelength between 400 and
2.498 nm) using a NIRS instrument fitted with a spinning cup holder (NIRSystem 6500; Perstorp Analytical Inc., Silver Spring, MD). Near infrared spectra (log 1/reflectance) were recorded for each 2-nm interval. Population characterization based on spectral variability of the samples was evaluated using the CENTER algorithm (Shenk and Westerhaus, 1991a,b) included in the WinISI II software package (version 1.5; Infrasoft International, Port Matilda, PA), with a maximum standardized Mahalanobis distance (H) from the average spectrum of 3.0. Low average neighbor spectra (NH) distance characterized the spectral database, leading to poor results in attempting a first calibration and equation using the whole data set of 1,281 original samples (data not shown). To avoid redundant samples and any over-fitting, which could cause misleading calibration and poor performance of prediction, spectra selection was performed by using the SELECT function of WinISI II software, a PCA analysis based on NH distance of the spectra population (Shenk and Westerhaus, 1991a,b). This procedure, with neighbor H set to 0.6, resulted in 301 sample spectra selected for the calibration data set as described above.

Mathematical treatments of spectral data were performed with WinISI II (version 1.5), exploring different math transformations for scatter correction and numbers of modified partial least squares (MPLS) terms in the models. The standard normal variate and detrending (SNV-D) method was applied for scatter correction (Barnes et al., 1989), together with first- or second-order derivative as mathematical treatment, to avoid noise not belonging to the sample that could bring misleading information to the spectra. The best predictive model was achieved using first-order derivative, with gaps and smoothing each 4 data points. The MPLS regression technique was used to develop chemometric models, using the full near infrared wavelength range (1,100–2,500 nm) and selecting a wavelength each 8 nm for a total of 173 wavelengths. An optimum number of MPLS terms was determined by maximizing coefficients of determination (R²) and minimizing standard errors of calibration (SEC; Shenk and Westerhaus, 1991a). Calibration performance was evaluated using the cross-validation technique (Shenk and Westerhaus, 1991b). The statistics used for the evaluation of the model were SEC and standard error of cross-validation (SECV) and coefficient of determination for the calibration (R²) and for the cross-validation (\(1 - VR\), where VR = variance ratio). The ratio of performance deviation (RPD), calculated as the ratio of the standard deviation of the original data to the SECV (Williams, 2004) and the range error ratio (RER), the ratio of the range of the original data to SECV, were also used to evaluate the calibration performance.

Fecal composition and in vitro fiber digestibility data are reported in Table 1. In the calibration data set (n = 301), not all samples were analyzed for each parameter. However, for fiber composition and 

Table 1. Summary statistics of fecal composition and 240-h in vitro digestibility for the samples used in the calibration data set

<table>
<thead>
<tr>
<th>Constituent (% of DM)</th>
<th>n (^2^)</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Range (^3^)</th>
<th>Mean</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>DM</td>
<td>228</td>
<td>10.02</td>
<td>21.99</td>
<td>11.97</td>
<td>14.90</td>
<td>2.02</td>
</tr>
<tr>
<td>CP</td>
<td>103</td>
<td>7.90</td>
<td>22.15</td>
<td>14.25</td>
<td>13.58</td>
<td>3.13</td>
</tr>
<tr>
<td>Starch</td>
<td>71</td>
<td>0.18</td>
<td>6.04</td>
<td>5.86</td>
<td>1.72</td>
<td>1.11</td>
</tr>
<tr>
<td>Ash</td>
<td>126</td>
<td>7.77</td>
<td>19.91</td>
<td>12.14</td>
<td>11.98</td>
<td>2.59</td>
</tr>
<tr>
<td>ADF</td>
<td>301</td>
<td>28.70</td>
<td>85.36</td>
<td>56.66</td>
<td>43.95</td>
<td>6.64</td>
</tr>
<tr>
<td>ADL</td>
<td>276</td>
<td>9.42</td>
<td>43.05</td>
<td>33.63</td>
<td>16.63</td>
<td>6.32</td>
</tr>
<tr>
<td>aNDFom</td>
<td>301</td>
<td>43.85</td>
<td>81.97</td>
<td>38.12</td>
<td>59.98</td>
<td>5.62</td>
</tr>
<tr>
<td>uNDF240 (% aNDFom)</td>
<td>300</td>
<td>27.36</td>
<td>59.35</td>
<td>31.99</td>
<td>43.88</td>
<td>6.27</td>
</tr>
<tr>
<td>IVNDFD240 (% aNDFom)</td>
<td>300</td>
<td>4.40</td>
<td>52.46</td>
<td>48.06</td>
<td>26.67</td>
<td>9.40</td>
</tr>
</tbody>
</table>

\(^1^\)aNDFom = amylase and sodium sulfite-treated NDF, corrected for ash residue; uNDF240 = undigested NDF estimated via 240-h in vitro fermentation; IVNDFD240 = in vitro aNDFom digestibility after 240 h of incubation.

\(^2^\)n = number of sample analyses.

\(^3^\)Range = maximum − minimum.
of the diet were immediately evident, except for CP, where the fecal concentration is affected by microbial protein and flaking cell excretion.

The number of MPLS terms, the number of samples used for NIRS equations, the calibration, and cross-validation statistics are reported in Table 2. The number of samples used in the calibration equations may be less than the total number of samples available, because outliers identified by the 2-tailed Student’s t-test, were not used to develop the calibration equations. The fraction of variance accounted for each MPLS calibration was relatively high for almost all components ($R^2 > 0.90$), except for DM ($R^2 = 0.77$) and starch ($R^2 = 0.66$). After cross-validation, the coefficient of determination ($1 − VR$) for almost all components was, on average, 0.06 points lower than the $R^2$ of the calibration, with the exception of DM and starch. For DM, the low correlation could be related to the fact that samples were dried before scanning, whereas for starch, the wide difference between $R^2$ and $1 − VR$ suggested that more samples are required to strengthen the calibration. However, the calibration for starch looked promising and as reported in previous work (Fredin et al., 2014), robust calibrations for fecal starch are already available, and many feed laboratories in the United States offer such NIRS analysis to professional customers.

In regards to fecal uNDF240, and considering the peaks’ correlation with SNV-D (standard normal variate and detrending) spectra, clear absorption was observed in the region 1,900 to 2,000 nm, at 2,450 nm, and in the regions 1,150 to 1,650 nm and 2,000 to 2,350 nm, as reported by Nousiainen et al. (2004). Compared with grass silage, the difference in the shape and amplitude of the peaks obtained in the present work can be related to higher levels of NDF and uNDF240 found in fecal samples. Even if comparable NIRS data for fecal samples (and especially for uNDF240 and IVNDFD240) are not available in the literature, the good correlation among peaks between spectra suggests that a good prediction equation could be developed. Williams and Sobering (1996) considered that RPD value should be at least 3 and RER at least 10. Looking at these accuracy indicators in Table 2, the RER values were, in all cases, $>10$ and, in some cases, $>20$ (aNDFom, ADF), and the RPD values were $>2.5$ for almost all components except DM (1.69), starch (1.35), and ADL (2.42), suggesting a proper calibration. Despite the limited number of samples, the relatively high $R^2$ and $1 − VR$ values for CP and ash could be related to the fact that CP is a nitrogen compound, chemically well defined, whereas ash represents the inorganic portion of the sample and is well predicted by NIRS as the difference between DM and OM. Compared with values for CP, the $R^2$ and $1 − VR$ values were lower for uNDF240 and IVNDFD240. As reported by Berzaghi et al. (1997) and shown by Nousiainen et al. (2004), these results were expected, because fiber fractions, in general, are not composed of a single defined compound but combine different chemical entities involving several regions of spectral absorptions, altering the real relationship between the spectra and reference method. In addition, the laboratory procedures for uNDF240, IVNDFD240, and aNDFom could have larger analytical error compared with that for CP and ash due to the complexity of such methods. Other authors have also noted that the main source of inaccuracy in the calibration process is sampling and internal laboratory error, in relationship to the range of the given nutrient

Table 2. Calibration and cross-validation statistics for near infrared spectroscopy analysis of fecal chemical composition and 240-h in vitro digestibility

<table>
<thead>
<tr>
<th>Constituent</th>
<th>n</th>
<th>MPLS terms</th>
<th>Mean</th>
<th>SD</th>
<th>SEC</th>
<th>$R^2$</th>
<th>SECV</th>
<th>1 − VR</th>
<th>RPD</th>
<th>RER</th>
</tr>
</thead>
<tbody>
<tr>
<td>DM</td>
<td>215</td>
<td>10</td>
<td>14.82</td>
<td>1.87</td>
<td>0.90</td>
<td>0.77</td>
<td>1.11</td>
<td>0.65</td>
<td>1.69</td>
<td>10.83</td>
</tr>
<tr>
<td>CP</td>
<td>92</td>
<td>9</td>
<td>13.30</td>
<td>2.90</td>
<td>0.74</td>
<td>0.93</td>
<td>1.08</td>
<td>0.86</td>
<td>2.68</td>
<td>13.20</td>
</tr>
<tr>
<td>Starch</td>
<td>65</td>
<td>6</td>
<td>1.48</td>
<td>0.74</td>
<td>0.43</td>
<td>0.66</td>
<td>0.55</td>
<td>0.51</td>
<td>1.35</td>
<td>10.74</td>
</tr>
<tr>
<td>Ash</td>
<td>116</td>
<td>9</td>
<td>11.81</td>
<td>2.49</td>
<td>0.75</td>
<td>0.91</td>
<td>0.98</td>
<td>0.84</td>
<td>2.53</td>
<td>12.34</td>
</tr>
<tr>
<td>aNDFom</td>
<td>287</td>
<td>14</td>
<td>59.93</td>
<td>5.29</td>
<td>1.48</td>
<td>0.92</td>
<td>1.89</td>
<td>0.87</td>
<td>2.80</td>
<td>20.19</td>
</tr>
<tr>
<td>ADF</td>
<td>285</td>
<td>6</td>
<td>43.73</td>
<td>6.06</td>
<td>1.85</td>
<td>0.91</td>
<td>2.07</td>
<td>0.88</td>
<td>2.93</td>
<td>27.41</td>
</tr>
<tr>
<td>ADL</td>
<td>249</td>
<td>13</td>
<td>15.61</td>
<td>4.96</td>
<td>1.34</td>
<td>0.93</td>
<td>2.05</td>
<td>0.85</td>
<td>2.42</td>
<td>16.39</td>
</tr>
<tr>
<td>uNDF240</td>
<td>277</td>
<td>14</td>
<td>44.04</td>
<td>5.76</td>
<td>1.65</td>
<td>0.92</td>
<td>2.24</td>
<td>0.86</td>
<td>2.57</td>
<td>14.30</td>
</tr>
<tr>
<td>IVNDFD240 (% of aNDFom)</td>
<td>276</td>
<td>12</td>
<td>26.09</td>
<td>8.69</td>
<td>2.68</td>
<td>0.90</td>
<td>3.43</td>
<td>0.85</td>
<td>2.53</td>
<td>14.00</td>
</tr>
</tbody>
</table>

aNDFom = amylase and sodium sulfite–treated NDF, corrected for ash residue; uNDF240 = undigested NDF estimated via 240-h in vitro fermentation; IVNDFD240 = in vitro aNDFom digestibility after 240-h incubation.

1n = number of samples used in the equation; MPLS terms = number of factors in modified partial least square equation; SEC = standard error of calibration.

2SECV = standard error of cross validation; 1 − VR = coefficient of determination of cross-validation, where VR = variance ratio; RPD = ratio of performance deviation (SD/SECV); RER = range error ratio (range/SECV).
(Mentink et al., 2006). Furthermore, the major problems in interpreting NIRS spectra, related to cell wall digestibility, are the high interference of residual water absorbance and the variable absorbance of C–H bonds in many spectral regions, which overlaps the absorbance of digestible components of aNDFom and other components (i.e., sugars, starch), as observed by Nousiainen et al. (2004). Despite the fact that the uNDF240 and IVDNDFD240 data are based on complex methods and time-consuming laboratory procedures, they are still reliable and accurate because measuring only the end-point of digestion could avoid problems associated with curve fitting.

Despite several factors that could influence the development of a NIRS calibration for uNDF240, IVNDFD240 and other constituents in fecal samples, the data obtained in the present work showed that uNDF240 should be considered a physical entity composed of several chemical entities that can be well predicted by NIRS. The use of uNDF240 values developed from NIRS calibrations, as an alternative to the traditional technique that requires ADL to estimate iNDF values, could improve the accuracy and use of feed in diet evaluation models for ruminants. Moreover, uNDF240 could be an excellent marker because it can be traced from the diet to the feces. The NIRS technique is well suited for digestibility studies and can be used from in vitro data to develop reliable calibrations and equations.

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