

# Article

# Assessment of macadamia kernel quality defects by means of near infrared spectroscopy (NIRS) and nuclear magnetic resonance (NMR)

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1	Assessment	of	macadamia	nut	quality	defects	by	means	of	near	infrared
2	spectroscopy	y (N	IRS) and nuc	lear 1	magnetic	resonan	ce (N	NMR)			

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#### 23 Abstract

24 Macadamia kernels are visually sorted based on the presence of quality defects 25 by specialized labors. However, this process is not as accurate as non-destructive methods such as near infrared spectroscopy (NIRS) and nuclear magnetic resonance 26 27 (NMR). Thus, NIRS and NMR in combination with chemometrics have become 28 established non-destructive method for rapid assessment of quality parameters in the 29 food and agricultural sectors. Therefore, the quality of macadamia nuts was assessed by 30 NIRS and NMR using chemometric tools such as PCA-LDA and GA-LDA to evaluate 31 kernel defects. Macadamia kernels were classified as: 1=good, marketable kernels 32 without defects; 2=kernels with discoloration; 3=immature kernels; 4=kernels affected 33 by mold; and 5=kernels with insect damage. Using NIRS, the GA-LDA resulted in an 34 accuracy and specificity of 97.8 % and 100 %, respectively, to classify good kernels. On 35 the other hand, PCA-LDA technique resulting in an accuracy higher than 68 % and 36 specificity of 97.2 % to classify immature kernels. For NMR, PCA-LDA resulted in an 37 accuracy higher than 83% and GA-LDA resulted in an accuracy of 100%, both to 38 classify kernels with insect damage. NIRS and NMR spectroscopy can be successfully 39 used to classify unshelled macadamia nuts based on the defects. However, NIRS out-40 performed NMR based on the higher accuracy results.

41

42 Keywords: *Macadamia integrifolia* Maiden & Betche, TD–NMR, PCA-LDA, GA43 LDA, chemometrics.

44

#### 46 **1. Introduction**

47 Macadamia (*Macadamia integrifolia* Maiden & Betche) nut growers are keen 48 to continuously improve nut quality as this is the main characteristic required by the 49 final consumers. Nogueira (2008) mentioned that the quality of macadamia fruit is 50 associated with favorable climatic conditions, planning and orchard management, 51 varieties, pest control, plant nutrition, harvest and post-harvest practices. All these 52 factors are decisive for macadamia development and nut quality.

53 According to O'Hare et al. (2004), the main defects that can be observed in 54 macadamia nuts are immaturity; small nuts; cracks in the shell that allow the occurrence 55 of biological and chemical contamination; lipids oxidation, which result in unpleasant 56 odor and taste; bruises, and high moisture. Guthrie et al. (2004) reported other defects 57 that may be considered, as such: fungal growth, decomposition, germination, and 58 discoloration of macadamia nuts. Therefore, sound and/or good macadamia nuts must 59 have light cream color, no signs of mold, decay, insect scars, blemishes, hollow centers, 60 dark centers, shriveling, off-odors, adhering shells, and loose of extraneous material 61 (Wall, 2013).

62 Macadamia industry has developed various parameters of quality standards. 63 The Southern African Macadamia Growers' Association (SAMAC) classifies 64 macadamia nuts into three classes: first grade, commercial grade, and local market. 65 These classes are established based on kernel color, flavor and odor, kernel dust, insect 66 infestation, foreign material. A limit of 1.5 % is used reject the nuts based on the presence of insect damage, discoloration, and immaturity (SAMAC, 2018). On the other 67 68 hand, the United Nations Economic Commission for Europe (UNECE) has a higher 69 tolerance (5 %) for the presence of these defects (UNECE, 2010).

The sorting process of macadamia kernels in the industry can be carried out manually (Piza, 2005) or electronically (France, 2007), but both present flaws, since manual sorting of defective kernels can decrease dramatically with the use of inadequate lighting and untrained personnel, and the electronic selection uses color to sort kernels, which may lead to improper selection, since immature kernels can only be identified based on the deformed, wrinkled, and shrunken kernel (SAMAC, 2018).

76 The increasing requirements of consumers, regulatory agencies, and 77 competitors have been an impulse for the development of more accurate quality 78 assessment techniques in the food industry. In this regard, near infrared spectroscopy 79 (NIRS) in combination with chemometric modelling have become an established 80 method for rapid assessment and non-destructive quality parameters in the food and 81 agricultural sectors (Abbott, 1999; Jensen et al., 2001), since it is fast, safe, relatively 82 inexpensive technique and provides automation of quality control processes in products 83 of agroindustry (Pasquini, 2003).

84 NIRS has been used to evaluate macadamia nut quality. Guthrie et al. (2004) 85 developed modified partial least squares regression (MPLS) models for oil content 86 determination in intact macadamia kernels with a root mean square error of calibration 87 (RMSEC) of 2.4 % and discriminated intact kernels with brown centers or rancidity 88 from each other and from sound kernels using PCA. Canneddu et al. (2016) developed 89 models for predicting peroxide value (PV) and acidity index (AI) using PLSR and 90 classification models to discriminate defects present on shelled macadamia nuts using 91 FT-NIR. The best model for PV prediction resulted in a coefficient of determination  $(R_p^2)$  of 0.72, and for AI prediction a SEP of 0.14 % and a  $R_p^2$  of 0.80. Adequate 92 93 classification models (93.2 %) for defects was possible using principal component 94 analysis linear discriminant analysis (PCA-LDA). Carvalho et al. (2017) classified

95 intact macadamia nuts according to cultivars using PCA-LDA and genetic algorithm 96 with linear discriminant analysis (GA-LDA), reporting an accuracy higher than 94.4 % 97 and a value of 82.7 % for sensitivity using GA-LDA, respectively. The better 98 performance of GA-LDA can be due to that GA algorithm selects several wavenumbers 99 in a single band, due to collinearity problems. Carvalho et al. (2019) evaluated the 100 oxidative stability in intact macadamia nuts during drying process and reported a SEP 101 of 0.55 meq.kg<sup>-1</sup> and R<sup>2</sup>c of 0.57 for PV prediction, and SEP of 0.14 % and R<sup>2</sup>c of 0.29 102 for AI prediction. These results demonstrate that NIRS can be used to assess the 103 oxidative stability of intact macadamia nuts.

104 Nuclear magnetic resonance (NMR) has also been stated as an alternative 105 method among non-destructive techniques to evaluate fruit quality (Abbott, 1999). TD-106 NMR has wide applications for qualitative and quantitative in food analysis (Conalgo, 107 1996). In this regard, Pedersen et al. (2000) combined low-field nuclear magnetic 108 resonance (LF-NMR) and PCA to classify rape and mustard seeds according on the 109 type of seed, obtaining two distinct groups and 100 % of explained variance. This 110 technique was also applied to evaluate the efficacy of hydrophobic coatings as a barrier 111 to the oxidation of macadamia nuts (Colzato et al., 2009).

Although some results can be found regarding the use of NIRS to assess macadamia quality defects (Canneddu et al., 2016), this study was performed evaluating the macadamia in nut not the kernel (unshelled), and no reports were found on using NMR to evaluate macadamia kernel defects. Therefore, the objective of this study was to develop NIRS and NMR calibration models to evaluate macadamia kernels based on the most common defects aiming to improve the quality control process in the macadamia industry.

#### 120 **2. Material and Methods**

#### 121 **2.1. Plant material**

122 Macadamia (Macadamia integrifolia Maiden & Betche) kernels were obtained 123 in a commercial orchard located in Dois Córregos, São Paulo, Brazil (22º 37' S, latitude, 124 48° 38' W, longitude, 753 m altitude) in 2017 harvest season. Nuts were harvested three 125 times during the season (April, June, and August) and kernels were visually sorted by 126 the industry personnel based on their quality attributes, as such: 1=good, marketable 127 kernels without defects; 2=kernels with discoloration; 3=immature kernels; 4=kernels 128 affected by mold; and 5=kernels with insect damage. (Figure 1). These quality attributes 129 represented the five studied classes (model). It is important to state that the nuts were 130 dried by the processing industry and used in the analyses without any previous 131 treatment.

#### 132 **2.2. NIR spectra acquisition**

133 On the surface of each macadamia kernel two Fourier Transformed (FT) NIR reflectance spectra  $(11,544 - 3,952 \text{ cm}^{-1}, \text{ nm}, \text{ resolution of } 16 \text{ cm}^{-1}, \text{ and } 64 \text{ scans})$  were 134 135 collected using a Bruker NIR spectrometer (Tango, Ettlingen, Germany) after 136 temperature stabilization at  $\sim 25^{\circ}$ C. The two replica spectra measured per nut were 137 averaged, so the model is made on a sample basis. Samples were collected in three 138 different harvests, where 20 nuts were sorted and used for spectra acquisition for each 139 defect class. This resulted in a total of 300 measured samples (20 nuts x 5 classes x 3 140 harvests).

#### 141 **2.3. Time domain (TD) NMR measurements**

142TD-NMR measurements of macadamia kernels (n=100) were carried out at 22143°C in a 0.27 T (11.3 MHz for <sup>1</sup>H) benchtop SLK200 Spinlock instrument (Spinlock144Magnetic Resonance Solutions, Cordoba, Argentina). The measurements were

145 performed using the standard CPMG sequence to obtain the exponential decay signal 146 that is governed by the transverse relaxation time (T2). The sequence used  $\pi/2$  and  $\pi$  of 147 11.6 and 19.6 µs, respectively, an echo time of 600 µs, 4 scans and 1500 echoes. 148 Samples harvested in June 2017 were used and for each defect class 20 nuts were sorted 149 and used for spectra acquisition, totaling 100 spectra. The mass of the samples ranged 150 from 14 to 24 g depending on the sample density. The samples were the same used to 151 collect the NIRS spectra, but the spectra were collected on different days.

#### **2.4. Chemometrics**

153 Data analysis of NIR and TD-NMR were performed within MATLAB R2014b 154 environment (MathWorks Inc., USA) using PLS Toolbox version 7.9.3 (Eigenvector Research Inc., USA) and lab-made routines. Three different pre-processing methods 155 156 were applied to test the averaged sample spectrum (average of 10 spectra per sample): 157 (1) only mean-centering; (2) standard normal variate (SNV) followed by meancentering; (3) Savitzky-Golay second derivative (window of 5 points, 2<sup>nd</sup> order 158 159 polynomial function) followed by mean-centering. The data was split into training (70 160 %, 210 samples), validation (15 %, 45 samples) and test (15 %, 45 samples) sets using 161 the Kennard-Stone sample selection algorithm (Kennard and Stone, 2012). The training 162 and validation sets were used for model construction and internal optimization, 163 respectively; while the test set was used to evaluate the final predictive performance of 164 the classification models built towards external samples.

Multivariate classification was performed by means of principal component analysis linear discriminant analysis (PCA-LDA) and genetic algorithm linear discriminant analysis (GA-LDA). PCA-LDA performs a feature extraction using principal component analysis (PCA) followed by a linear discriminant classifier (LDA) (Morais and Lima, 2018) For this, PCA is applied to the pre-processed data reducing the original number of variables (i.e., wavelengths) to a few number of principal
components (PCs) accounting for the majority of the original data variance. Each PC is
composed by scores and loadings, where the first represents the variance between the
samples and the latter the variance on wavelength direction (Bro and Smilde, 2014).
LDA is applied to the PCA scores in a non-Bayesian form as follows (Dixon and
Brereton, 2009; Wu et al, 1996).

176 
$$L(\mathbf{x}_i) = (\mathbf{x}_i - \bar{\mathbf{x}}_k)^{\mathrm{T}} \mathbf{C}_{\text{pooled}}^{-1} (\mathbf{x}_i - \bar{\mathbf{x}}_k)$$
(1)

177 where  $L(\mathbf{x}_i)$  represents the LDA classification scores for sample *i*;  $\mathbf{x}_i$  is the input vector

178 (i.e., the PCA scores) for sample  $i; \bar{\mathbf{x}}_k$  is the average vector of class  $k; \mathbf{C}_{pooled}$  is pooled

179 covariance matrix; and **T** represents the matrix transpose operation.

GA-LDA is feature selection technique followed by an LDA classifier. Initially, a genetic algorithm (GA) is applied to reduce to the spectral data into a few number of variables based on an evolutionary process (Bro and Smilde, 2014); then LDA is applied to these variables according to Eq. 1. These variables are in the same scale of the original spectral data and are selected according to the lowest risk of miss classification *G*. *G* is calculated in the validation set as (Carvalho et al. 2017).

186 
$$G = \frac{1}{N_{\rm v}} \sum_{n=1}^{N_{\rm v}} g_n \tag{2}$$

187 where  $N_v$  is the number of validation samples and  $g_n$  is defined as:

188 
$$g_n = \frac{r^2(x_n \cdot m_{I(n)})}{\min_{I(m) \neq I(n)} r^2(x_n \cdot m_{I(m)})}$$
(3)

189 in which the numerator is the squared Mahalanobis distance between sample  $x_n$  (of

190 class index I(n) and the mean  $m_{I(n)}$  of its true class; and the denominator represents

191 the squared Mahalanobis distance between sample  $x_n$  and the mean  $m_{I(m)}$  of the

192 closest wrong class. GA was performed through 100 generations, having 200193 chromosomes each. Cross-over and mutation probabilities were set at 60% and 1%,

194 respectively. The algorithm was repeated three times and the best result was chosen.

195

#### 196 **2.5. Figures of merit**

197 The classification performance of each algorithm was evaluated according to 198 the quality parameters of accuracy (total number of samples correctly classified 199 considering true and false negatives), sensitivity (proportion of positives correctly 200 identified) and specificity (proportion of negatives correctly identified). These 201 parameters are calculated as follows (Morais and Lima, 2017):

202 
$$\operatorname{Accuracy}(\%) = \frac{\mathrm{TP} + \mathrm{TN}}{\mathrm{TP} + \mathrm{FP} + \mathrm{TN} + \mathrm{FN}} \times 100$$
(4)

203 Sensitivity (%) = 
$$\frac{TP}{TP+FN} \times 100$$
 (5)

204 Specificity (%) = 
$$\frac{\text{TN}}{\text{TN+FP}} \times 100$$
 (6)

205 where TP stands for true positives; TN for true negatives; FP for false positives; and FN

for false negatives.

207

#### 208 **3. Results and Discussion**

#### **3.1. NIR spectra**

The raw FT-NIR spectra obtained from all macadamia kernels and the average spectra from each quality attribute class can be seen in Figure 2. It was not possible to observe spectral differences between the quality attributes when all macadamia kernels were assessed (Figure 2A). On the other hand, the mean spectra were quite different for each defect category (Figure 2B), especially at the wavelength 1,900 nm to 2,500 nm.

215 The FT-NIR spectra presented absorption bands at 1,200 nm, which are related 216 to CH stretch second overtone (Cozzolino et al., 2005), while those at 1,700 - 1,800 nm 217 are associated to the first overtones of CH stretching vibrations of -CH<sub>3</sub>, -CH<sub>2</sub>- and -218 HC=CH (Armenta and La Guardia, 2007). Absorption bands at 1,350 - 1,600 nm and 219 1,950 nm and 2,100 nm are related to the presence of glucose, sucrose, and fructose 220 (Lanza and Li, 1984) and immature kernels have higher sucrose and reducing sugar 221 contents than fully mature kernels (Wall, 2013). In Figure 2B can be seen that at 1,350 222 -1,600 nm the immature kernels exhibit a higher absorption intensity, since maturity is 223 inversely related to sugar content (Ripperton et al., 1938).

The wavelength region situated at 2,200 – 2,500 nm is mainly related to the oxidation and hydrolytic degradation of lipids (Cozzolino et al. 2005). It is possible to observe that the immature kernels, classified as kernel which is misshapen, abnormally small or partially aborted, including shriveled and shrunken kernels (SAMAC, 2016) present a lower absorption band (2,200 nm - 2,500 nm) (Figure 2B). This result might be due to the fact that maturity is correlated with oil content (Cavaletto, 1985), consequently with less lipid degradation.

*3.1.1. Model development* 

235 Regarding pre-processing, SNV lead to best results using PCA-LDA, resulting 236 in an accuracy of 68 % and a specificity of 97 % for immature kernels (Table 1). The 237 accuracy shows the proportion of samples correctly grouped, while specificity 238 represents the probability of a sample without the desired characteristic to be given a 239 negative test result (Amodio et al., 2017). However, the sensitivity presented low 240 values (67 %), and this parameter describes the model ability to correctly recognize 241 samples belonging to a class (Ballabio and Consonni, 2013). For example, if none of 242 the marketable kernels were classified as other class (FN is equal to zero), the 243 sensitivity for the marketable kernels class would have been equal to 100 %.

Cannedu et al. (2016) classified marketable macadamia kernels in relation to non-marketable kernels using PLS-DA and reported percentages of 88 % for calibration and 87 % for prediction. These results were inferior than what we obtained, probably because we used more samples (n = 300) than Cannedu et al. (2016) (n = 100). Therefore, the inclusion of more data into the dataset improved the robustness and increase the classification accuracy.

250 Marketable kernels and kernels with defects (immature, insect damage, mold, 251 and discoloration) could be discriminated from each other using GA-LDA (Figure 3). 252 The accuracy and specificity of GA-LDA for marketable kernels achieved a value of 253 97.8 % and 100 %, respectively (Table 2).

To perform the GA-LDA, some of the wavelengths were selected (Table 3). This selection was based on compounds of particular interest, e.g., 1,020 nm and 1,173 nm, representing the C–H groups from lipids; 1,485 nm and 1,789 nm, related to the first overtone of stretching and anti-symmetric O–H bond and second overtone of stretching O–H bend, respectively. Absorption bands at the wavelength near 1,450 and 1,940 nm are related to the presence of water in foods (Moscetti et al., 2014) and this explains why the wavelengths 1,485 nm, 1,975 nm and 1,987 nm were selected by GA.

It is possible to observe that the kernels with discoloration had a higher moisture content than the others (Figure 2B), and these moisture contents correspond to water activities (aw) greater than 0.8 at which browning reaction rates are high (Wall, 2013), and maintaining nuts-in-shell at high moisture content can cause discoloration (Walton et al., 2013).

266 **3.2. TD-NMR** 

The typical curves of the CPMG decays for the different defects found in macadamia kernels can be seen in Figure 4. It can be observed that kernels with insect damage presented a faster settling time compared to the others, whereas the kernels with presence of fungi (moldy) showed the slower signal decay (Figure 4).

271 The intensity of the TD-NMR signals from relaxation (our case) and diffusion 272 measurements is related to the water content related to water status, water 273 compartmentalization and molecular mobility in the food sample (Kirtil et al., 2017). In 274 order to evaluate the influence of the water content on the nutrient content of the food, it 275 is important to note that there are variations in the moisture content of the kernels, since 276 these moisture contents correspond to water activities at which microbial growth rates 277 are high (Wall, 2013). This explains the fact that moldy kernels have a higher moisture 278 content.

In Figure 5 it is possible to observe that there was not a clear separation between the defect classes. However, in Figure 5A there was a tendency of separation between the good and immature kernels. Probably because there are differences in the decay time between these classes (Figure 4), with showed that the most rapid decay is due to solid components, mainly composed of proteins and carbohydrates (Prestes et al., 2007) and immature kernels present a higher carbohydrate concentration, represented by sucrose and fructose higher than mature kernels (Wall, 2013).

286 *3.2.1. Model development* 

The best TD-NMR classification models were obtained using the PCA-LDA and GA-LDA without pre-processing the signals (Table 4). Using PCA-LDA, it was possible to achieve 86 % accuracy for the training set and 83.3 % for the validation set to classify kernels with insect damage. On the other hand, the GA-LDA analysis obtained 64 % for the calibration set and 100 % for the validation set, allowing the use of this model to classify kernels with insect damages.

TD-NMR has been used to classify other oleaginous produces including nuts. Di Caro et al. (2017) studying not damaged and moldy hazelnuts kernels highlighted that NMR might be used to discriminate oils extracted from both kernel classes. Di Caro (2018) also reported that using NMR was possible to obtain values of 97 % for sensitivity and 81 % for specificity to classify in-shell damaged hazelnuts. Therefore, NMR might be a useful analytical tool for quality control in nut industry.

299 **3.3. NIRS versus TD-NMR** 

The results obtained from both techniques for the development of the classification models for macadamia kernels quality defects can be seen in Table 1, 2, and 4. Overall, the NIRS showed better classification capability as higher values of accuracy were obtained using GA-LDA models. The lower performance of the classification models developed using the TD-NMR signals might be related to the number of samples, as just the kernels harvested in June 2017 were used. 306 NIRS and TD-NMR present many similarities as they are fast non-destructive 307 analytical methods, do not need sophisticated sample preparation, and the results can be 308 collected, processed, and stored directly in a microcomputer (Colnago, 1996; Pasquini, 309 2003). However, when it comes to NMR spectroscopy, high cost is normally considered 310 as one of the most serious drawbacks and this technique requires special skills to 311 interpret the spectra acquisition (Xu et al., 2015). Another limitation of NMR 312 spectroscopy is the insensitivity to minor fat component detection (Kucha et al., 2018). 313 These suggest that, due the fact that NIRS is useful for detecting components with up to 314 0.1 % concentration (Xu et al., 2015) and NMR presents lower sensitivity, NIRS models 315 presented more satisfactory results. 316 317 4. Conclusions 318 NIRS and TD-NMR combined with chemometric methods proved to be 319 powerful tools to classify macadamia kernels based on their quality defects. However, 320 NIRS out-performed TD-NMR based on the higher accuracy results. 321 NIRS and TD-NMR spectroscopy can be successfully used to evaluate the 322 quality of unshelled macadamia nuts and have potential to improve the existing 323 postharvest techniques used in the macadamia industry. 324 325 Acknowledgment 326 This study was financed in part by the Coordenação de Aperfeiçoamento de 327 Pessoal de Nível Superior – Brasil (CAPES) – Finance Code 001. 328 329 References 330 Abbott, J.A. (1999). Quality measurement of fruit and vegetables. Postharvest Biology 331 and Technology, 15, 207-223.

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## 434 **Tables**

- 435 **Table 1**. Values of accuracy, sensitivity and specificity to classify macadamia kernels
- 436 based on quality defects using PCA-LDA and NIRS.

Classes		1	2	3	4	5
Pre-Processing						
Raw	AC(%)	88.9	84.4	75.6	82.2	75.6
	SENS(%)	88.9	66.7	44.4	44.4	22.2
	SPEC(%)	88.9	88.9	83.3	91.7	88.9
SNV	AC(%)	80.0	68.9	88.9	75.6	75.6
	SENS(%)	66.7	55.6	55.6	11.1	33.3
	SPEC(%)	83.3	72.2	97.2	91.7	86.1
2 <sup>nd</sup> Derivative	AC(%)	82.2	73.3	86.7	88.9	75.6
	SENS(%)	66.7	44.4	77.8	66.7	11.1
	SPEC(%)	86.1	80.6	88.9	94.4	91.7

437 SNV= standard normal variate; AC= accuracy; SENS= sensitivity; SPEC= specificity.

438 1=good, marketable kernels without defects; 2=kernels with discoloration; 3=immature

439 kernels; 4=kernels affected by mold; and 5=kernels with insect damage.

**Table 2**. Values of accuracy, sensitivity and specificity to classify macadamia kernels

444	based on quality defects using GA-LDA and NIRS.
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Classes		1	2	3	4	5
Pre-Processing						
Raw	AC(%)	86.7	82.2	86.7	86.7	82.2
	SENS(%)	66.7	66.7	55.6	66.7	55.6
	SPEC(%)	91.7	86.1	94.4	91.7	88.9
SNV	AC(%)	97.8	84.4	88.9	91.1	84.4
	SENS(%)	88.9	88.9	55.6	77.8	55.6
	SPEC(%)	100	83.3	97.2	94.4	91.7
2 <sup>nd</sup> Derivative	AC(%)	91.1	75.6	84.4	86.7	68.9
	SENS(%)	66.7	44.4	44.4	55.6	55.6
	SPEC(%)	97.2	83.3	94.4	94.4	72.2

445 SNV= standard normal variate; AC= accuracy; SENS= sensitivity; SPEC= specificity.

446 1=good, marketable kernels without defects; 2=kernels with discoloration; 3=immature

447 kernels; 4=kernels affected by mold; and 5=kernels with insect damage.

- **Table 3.** Selected variables for GA-LDA to classify macadamia kernels using different
- 452 pre-processing.

Pre-processing	Selected variables (nm)
Raw	882; 886; 946; 990; 1171; 1395; 1429; 1511; 1622; 1664; 1942;
	1979; 2075; 2187; 2260; 2328
SNV	866; 1020; 1173; 1280; 1485; 1578; 1789; 1975; 1987; 2083;
	2170; 2277; 2300; 2388; 2451
2 <sup>nd</sup> Derivative	894; 898; 1078; 1251; 1335; 1436; 1488; 1952; 1964; 2126;
	2328; 2356

453 SNV=standard normal variate

Classes		1	2	3	4	5
Pre-Processing						
	PCA-LDA					
	Training (%)	64.3	35.7	42.9	85.7	64.3
Nil	Validation (%)	16.7	33.3	16.7	66.7	83.3
	GA-LDA					
	Training (%)	64.3	50.0	35.7	64.3	50.0
	Validation (%)	66.7	16.7	66.7	66.7	100

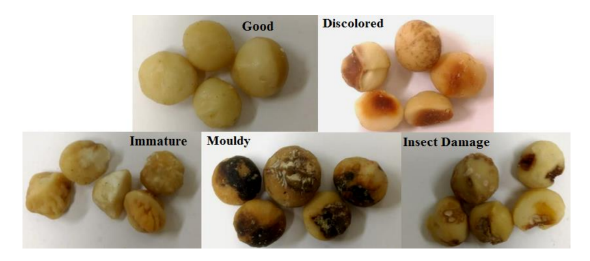
457 using PCA-LDA, GA-LDA and TD-NMR spectroscopy.

458 1=good, marketable kernels without defects; 2=kernels with discoloration; 3=immature

459 kernels; 4=kernels affected by mold; and 5=kernels with insect damage.

460

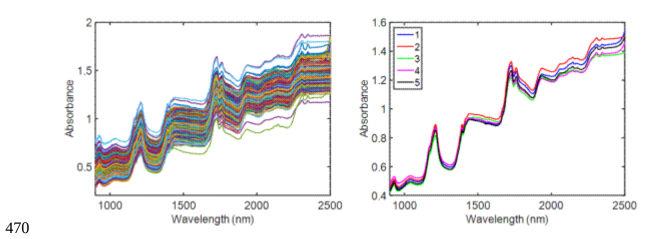
## **Figures**



464 Figure 1. Macadamia kernels quality defects: 1=good, marketable kernels without

465 defects; 2=kernels with discoloration; 3=immature kernels; 4=kernels affected by mold;

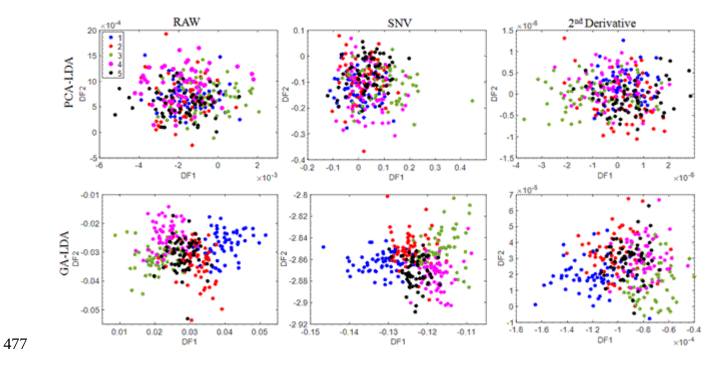
- 466 and 5=kernels with insect damage.



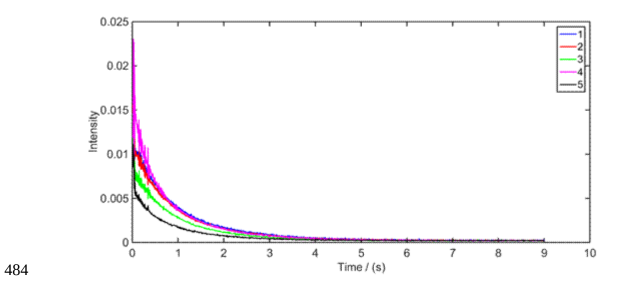
471 Figure 2. Raw NIR spectra (a) and average NIR spectra (b) of macadamia kernels.

472 1=good, marketable kernels without defects; 2=kernels with discoloration; 3=immature

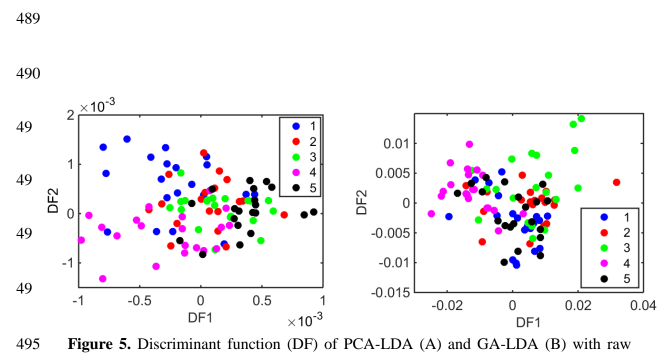
- 473 kernels; 4=kernels affected by mold; and 5=kernels with insect damage.
- 474



478 Figure 3. Discriminant function (DF) plot of PCA-LDA and GA-LDA with raw NIR
479 spectra of macadamia kernels, SNV and 2<sup>nd</sup> derivative Savitzky-Golay. 1=good,
480 marketable kernels without defects; 2=kernels with discoloration; 3=immature kernels;
481 4=kernels affected by mold; and 5=kernels with insect damage.



485 Figure 4. Typical CPMG decay curves of macadamia kernels with different quality
486 defects. 1=good, marketable kernels without defects; 2=kernels with discoloration;
487 3=immature kernels; 4=kernels affected by mold; and 5=kernels with insect damage.



TD-NMR spectra of macadamia kernels. 1=good, marketable kernels without defects;
2=kernels with discoloration; 3=immature kernels; 4=kernels affected by mold; and
5=kernels with insect damage.