

**Investigation of the Potential Utility of Single-Bounce
 Attenuated Total Reflectance Fourier Transform Infrared
 Spectroscopy in the Analysis of Distilled Liquors and Wines**

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A new Fourier transform infrared (FTIR) spectroscopic method based on single-bounce attenuated total reflectance (SB-ATR) spectroscopy was developed for the analysis of distilled liquors and wines. For distilled liquors, a partial least-squares (PLS) calibration was developed for alcohol determination based on the SB-ATR/FTIR spectra of mixtures of ethanol and distilled water. An independent set of 12 different distilled liquor samples was predicted from the PLS calibration, and a standard deviation of the differences for accuracy (SDD_a) between actual and predicted values of 0.142% (v/v) was obtained. The potential utility of SB-ATR/FTIR spectroscopy for the analysis of wines was initially evaluated on the basis of a comparison of PLS calibrations of preanalyzed wine samples ($n = 28$) with Fourier transform near-infrared (FT-NIR) spectroscopy and FTIR spectroscopy using a flow-through transmission cell. PLS calibrations for alcohol, total reducing sugars, total acidity, and pH were evaluated, and for SB-ATR/FTIR spectroscopy, the SDD_a for the leave-one-out cross-validation statistics were of the order of 0.100% (v/v), 0.707 g L⁻¹, 0.189 g L⁻¹ (H₂SO₄), and 0.230, respectively. Overall, the SB-ATR/FTIR results were better than those obtained using FT-NIR spectroscopy and comparable to those obtained with transmission FTIR spectroscopy. A PLS calibration based on preanalyzed wine samples ($n = 72$) for the prediction of 11 different components and parameters in wines by SB-ATR/FTIR spectroscopy was subsequently developed and validated using an independent sample set ($n = 77$). Good coefficients of correlation between the reference and predicted values for the validation set were obtained for most of the components and parameters except citric acid, volatile acids, and total SO₂. The results of this study demonstrate the suitability of SB-ATR/FTIR spectroscopy for the routine analysis of distilled liquors and wines.

KEYWORDS: Wine; SB-ATR; FTIR; distilled liquors; PLS

INTRODUCTION

Spirits are largely alcohol/water mixtures with only minor levels of added or developed constituents present. Wines, on the other hand, can have a complex provenance, and their quality depends on both volatile (e.g., ethanol and aromatic compounds) and nonvolatile (e.g., sugars, organic acids, tannins, nitrogen compounds, and mineral matter) constituents (1, 2). As a consequence, whereas the analysis for alcohol in spirits and wines is routine for industrial quality control and government regulatory purposes, more detailed analysis for other constituents in wines is also commonplace (3, 4). Such analyses tend to be tedious, and the availability of a simple and rapid means of obtaining most of the relevant information using a single method of analysis would clearly be desirable.

Near-infrared (NIR) spectroscopy might be considered a likely candidate to meet this demand as it is a well-established

technique for multicomponent analysis of agricultural and food products (5). Its utility for the analysis of alcohol in wines was initially demonstrated using filter-based NIR reflectance spectrometers, employing wavelengths of 2270, 2230, 2180, and 1778 nm (6). Since then, NIR spectroscopic methods have also been developed for the analysis of alcohol in spirits using transmission measurements (7, 8) and flow injection analysis (9), for monitoring alcohol and sugars during wine fermentation (3), and for analyzing alcohol, sugars, and glycerol in botrytized-grape sweet wines using filter-based reflectance spectrometers (10). Recently, an FT-NIR transmission method was developed for the analysis of methanol in grape-derived distillation products (11) and a reflectance NIR method was developed for the analysis of 15 parameters in wines employing partial least-squares (PLS) regression (12). Overall, although NIR spectroscopy is an adequate analytical tool for predicting the alcohol content in wines and spirits, it is limited in its ability to analyze other wine components. For example, although NIR methods for the determination of sugar content have been developed for

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sweet wines (10), the major NIR absorption bands of sugars are highly overlapped by the OH stretching and deformation combination bands of water, making the measurement of sugars imprecise below 30 g L⁻¹ (13).

Although mid-IR spectroscopy has generally found much less practical application than NIR spectroscopy in the analysis of agricultural and food products, it has been successfully employed in the analysis of alcohol in alcoholic beverages (14, 15) and has recently been shown to be a suitable technique for the analysis of multiple components in wines, including tartaric acid, malic acid, lactic acid, total acidity, pH, volatile acidity, reducing sugars, fructose, glucose, and glycerol (16–18). However, at present, Fourier transform infrared (FTIR) wine analysis is basically restricted to proprietary commercial instrumentation dedicated to this application but based on the design of FTIR milk analyzers. The sample-handling accessory integrated into these systems incorporates a temperature-controlled flow-through transmission cell equipped with CaF₂ windows and having a fixed path length of ~50 μm. Despite this short path length, which is required to limit the overwhelming IR absorptions of water (18), products containing high levels of alcohol and/or sugar would require dilution prior to analysis in order to keep the major absorption bands of these components on scale. The present work was undertaken to develop FTIR analytical methodology that utilizes a more versatile single-bounce attenuated total reflectance (SB-ATR) sample-handling accessory and a standard benchtop FTIR spectrometer, thereby reducing the cost of implementing FTIR wine analysis. The advantages of the recently developed SB-ATR accessories (19) were exploited in previous work in our laboratory for the quantitative analysis of sugars during lactose hydrolysis in milk (20). These include higher energy throughput (19) and more precise temperature control by comparison with the multiple-reflection devices traditionally employed for ATR measurements. Furthermore, a ZnSe SB-ATR accessory provides access to much more spectral information than the transmission cell employed in commercial FTIR wine analyzers owing to its very short path length, as well as the much lower transmission cutoff of ZnSe compared to that of CaF₂. On the basis of these considerations, the present study was undertaken to evaluate the utility of SB-ATR/FTIR spectroscopy for the analysis of distilled liquors and wines. In the first part of the study, alcohol determination in spirits, which are relatively simple matrices, was assessed. Subsequently, a comparative evaluation of SB-ATR/FTIR spectroscopy relative to FT-NIR and FTIR transmission methods for the analysis of wines was carried out. Finally, the simultaneous determination of 11 constituents in wines by SB-ATR/FTIR spectroscopy was investigated.

MATERIALS AND METHODS

Samples, Chemicals, and Reagents. Anhydrous alcohol (ethanol) was obtained from Commercial Alcohols Inc. (Brampton, ON, Canada), and 12 distilled liquor samples (rum, gin, vodka, brandy, scotch, and whiskey) were obtained from local liquor stores. A total of 177 wine samples were obtained courtesy of the central laboratory of the Société des alcools du Québec (SAQ), all of which had been analyzed for alcohol content by distillation and densitometry, for pH by potentiometry, and for total reducing sugars, total acidity, volatile acidity, and total SO₂ by colorimetry using an automated system (Skalar Analytical Inc., Tinststraat, The Netherlands). The products included red and white, dry, semisweet, and sweet wines from a wide variety of countries including Italy, France, Spain, the United States, South Africa, Germany, Canada, and Australia. All reagents were obtained from Fisher Scientific (Montreal, PQ, Canada) and were of analytical grade.

Instrumentation. All quantitative IR analyses were carried out using an ABB Bomem (Québec City, PQ, Canada) MB-150 dual-range (NIR/

mid-IR) FTIR spectrometer equipped with a deuterated triglycine sulfate detector. The spectrometer was purged with dry air from a Balston air drier (Lexington, MA) to avoid spectral contributions from atmospheric water vapor and carbon dioxide. SB-ATR spectra were recorded with the use of a Harrick (Ossining, NY) ZnSe accessory equipped with a temperature controller (Watlow System Integrators, Decorah, IA). During sample analysis, the SB-ATR crystal was covered with a glass microscope slide to avoid any alcohol evaporation during spectral recording, and the sample temperature was maintained at 30 ± 0.1 °C to minimize spectral changes due to temperature variation. For purposes of comparison, FT-NIR and FTIR analyses were carried out with a 1000-μm quartz cell and a 50-μm CaF₂ cell, respectively, both thermostated at 30 ± 0.1 °C using an Omega temperature controller (Omega Engineering, Stamford, CT). Spectral collection parameters for FT-NIR, transmission FTIR, and SB-ATR/FTIR analyses were 16, 32, and 32 co-added scans at resolutions of 16, 8, and 8 cm⁻¹, respectively. The spectra of all samples were ratioed against a background single-beam spectrum recorded immediately prior to collection of the spectrum of the sample.

HPLC analysis of wines for glycerol, fructose, and organic acids (citric acid, malic acid, tartaric acid, and lactic acid) was carried out employing a Hewlett-Packard 1050 HPLC system, equipped with an ultraviolet detector set to 192 nm. Separation was carried out isocratically on a 7.8 mm × 300 mm Rezex ROA-Organic Acid column manufactured by Phenomenex Inc. (Torrance, CA) using 0.0025 N sulfuric acid at a flow rate of 0.5 mL min⁻¹ and a column temperature of 52 ± 0.1 °C.

Calibration Development and Validation. For the determination of alcohol in spirits, calibration standards were prepared by blending anhydrous ethanol with distilled water to cover a range of 25–75% (v/v) alcohol, and the validation set comprised 12 distilled liquor samples. Both standards and validation samples were analyzed using an Anton Paar density meter (Graz, Austria) according to the Association of Official Analytical Chemists' method 982.10 (21). For wines, an initial set of 28 samples that had been analyzed for alcohol content, pH, total acidity, and total reducing sugars by standard methods was used for comparing the utility of the SB-ATR method to more conventional FT-NIR and FTIR transmission methods. A larger set of 149 wines was then used to assess the feasibility of employing FTIR/SB-ATR spectroscopy for the simultaneous determination of 11 components (alcohol, total reducing sugars, fructose, total acidity, volatile acidity, tartaric acid, malic acid, citric acid, lactic acid, glycerol, and SO₂), with 72 of the wines used for calibration and 77 serving as validation samples.

All calibrations were developed by applying PLS regression (22) using Omnic TQ Analyst software (Nicolet Instrument, Madison, WI). The correlation, variance, and pure component spectra generated from the PLS software package were used to select regions in developing and optimizing the calibration model for each component. The optimal number of factors used for each component was determined from the minimum point on the predicted residual error sum of squares (PRESS) plot. The overall suitability of the calibration model ultimately developed was assessed by calculating the mean difference for accuracy (MD_a), standard deviation of the differences for accuracy (SDD_a), and standard error of cross-validation (SECV) between the leave-one-out cross-validation predictions and values obtained by the reference method(s). For validating the FTIR/SB-ATR method employing 77 preanalyzed wine samples, the performance of the method was evaluated by calculating the MD_a, SDD_a, and standard error of prediction (SEP) between the predicted and reference values.

RESULTS AND DISCUSSION

Distilled Liquor Analysis. Although there are many ways of measuring alcohol in distilled liquors, the most commonly used official methods are those involving physical measurements (21, 23–25). NIR (7–9) and mid-IR (14, 15) transmission methods have also been investigated and developed, with NIR methods dominating. Owing to the simplicity of the distilled liquor matrix, the NIR spectra of distilled liquors exhibit some

SB-ATR FTIR Analysis of Distilled Liquors and Wines

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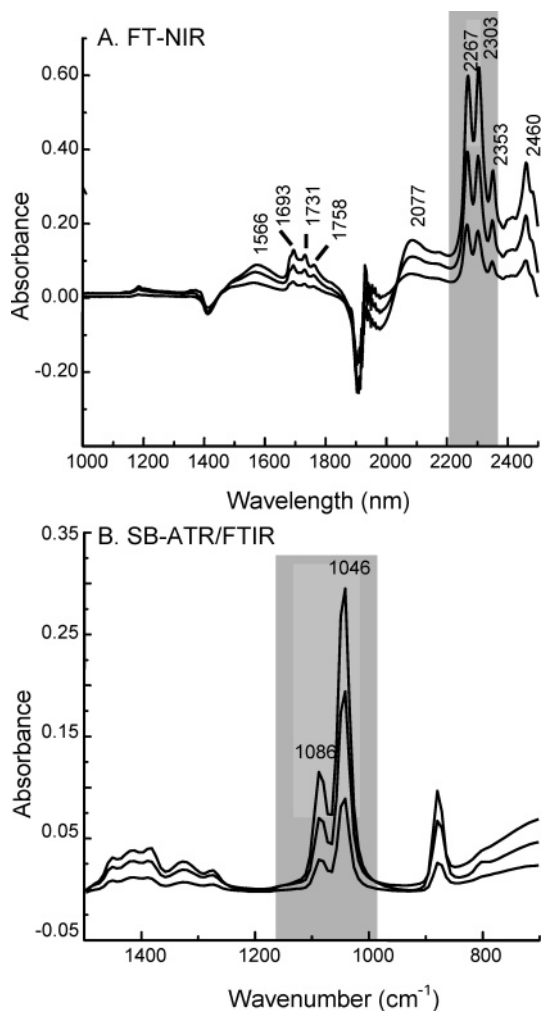


Figure 1. FT-NIR (A) and SB-ATR/FTIR (B) spectra of 20, 40, and 60% (v/v) solutions of ethanol after subtraction of water absorptions. The shaded area represents the spectral region ultimately used for PLS calibration.

198 fairly well-defined combination and first overtone bands of the
 199 O–H, C–H, and C–O stretching and bending vibrational
 200 frequencies of ethanol (8), whereas the mid-IR spectra exhibit
 201 two strong bands at 1046 and 1086 cm^{-1} due to the fundamental
 202 C–O stretching vibrations (**Figure 1**). Because transmission FT-
 203 NIR spectroscopy has been shown to be a suitable technique
 204 for the determination of alcohol in distilled liquors (7, 8), it
 205 was used as a basis of comparison in the evaluation of an SB-
 206 ATR/FTIR method. The spectra of ethanol/distilled water
 207 mixtures were used to develop PLS calibrations for the
 208 transmission FT-NIR and SB-ATR/FTIR procedures, and the
 209 spectral regions ultimately used to derive the PLS calibrations
 210 are shown in **Figure 1**. The calibrations were assessed by leave-
 211 one-out cross-validation, and linear regression of the cross-
 212 validation predictions versus the actual values yielded an R value
 213 of 0.999 for both methods with an SECV of 0.065 and 0.080%
 214 (v/v) for the FT-NIR and SB-ATR/FTIR methods, respectively.

215 The PLS calibrations were used to evaluate the alcohol
 216 content of 12 distilled liquors representing a variety of products.
 217 The predicted alcohol contents were compared to the results
 218 obtained using the Anton Paar density meter giving MD_a and
 219 SDD_a values of 0.03 and 0.163% (v/v) for FT-NIR and 0.02
 220 and 0.142% (v/v) for SB-ATR/FTIR; however, regression was
 221 not performed as there was inadequate variability in the alcohol
 222 content [ranging from 40 to 41% (v/v)] to produce meaningful
 223 regression data. The MD_a and SDD_a data indicate that the

Table 1. Regions Employed in FT-NIR, Transmission FTIR, and SB-ATR/FTIR PLS Regressions for a Comparative Study Employing 28 Wine Samples

component	FT-NIR regions (nm)	FTIR regions (cm^{-1})	
		transmission	SB-ATR
alcohol	2368–2207	1000–1500	850–1500
reducing sugar	1851–1666	1108–1500	956.7–1500 3000–2800
total acidity	2368–2207	1108.3–1500 1700–1800	1108.3–1500 1700–1800
pH	1851–1389	1000–1500	956.7–1500 3000–2800

Table 2. Cross-Validation Results for PLS Calibration Models Developed for the Determination of Alcohol, Reducing Sugar, Total Acidity, and pH by FT-NIR, FTIR, and SB-ATR/FTIR Spectroscopy Based on a Calibration Set of 28 Wine Samples, Including Linear Regression Equations and Values of R , SECV, MD_a , and SDD_a

component	method	equation	R	SECV	MD_a	SDD_a
alcohol [% (v/v)]	FT-NIR	$Y = 0.231 + 0.982X$	0.991	0.110	0.00	0.109
	FTIR	$Y = 0.059 + 0.996X$	0.991	0.111	0.01	0.109
	SB-ATR	$Y = 0.061 + 0.995X$	0.993	0.102	0.00	0.100
reducing sugar (g L^{-1})	FT-NIR	$Y = 1.479 + 0.341X$	0.500	0.764	−0.04	1.123
	FTIR	$Y = 0.946 + 0.584X$	0.735	0.697	−0.01	0.863
	SB-ATR	$Y = 0.605 + 0.711X$	0.837	0.613	−0.06	0.707
total acidity ($\text{g L}^{-1} \text{H}_2\text{SO}_4$)	FT-NIR	$Y = 2.874 + 0.152X$	0.270	0.167	0.00	0.304
	FTIR	$Y = 0.453 + 0.863X$	0.882	0.142	−0.01	0.146
	SB-ATR	$Y = 1.164 + 0.656X$	0.787	0.156	0.00	0.189
pH	FT-NIR	$Y = 2.107 + 0.390X$	0.548	0.202	0.01	0.283
	FTIR	$Y = 1.213 + 0.654X$	0.706	0.222	0.02	0.246
	SB-ATR	$Y = 1.416 + 0.582X$	0.713	0.190	−0.01	0.230

224 alcohol content of the distilled liquor products investigated could
 225 be predicted to within 0.17% (v/v) of the density meter values,
 226 with no significant bias indicated between the reference and IR
 227 methods, and that the FT-NIR and SB-ATR/FTIR methods
 228 performed comparably.

229 These results indicate that the SB-ATR/FTIR method is
 230 readily able to quantify alcohol content in spirits to a similar,
 231 if not higher, degree of accuracy as the FT-NIR procedure. The
 232 results are also comparable to those reported in publications on
 233 both NIR (7–9) and mid-IR (14, 15) methods for the analysis
 234 of distilled liquors. With respect to the previous mid-IR methods,
 235 which were based on the use of transmission cells, the SB-
 236 ATR procedure has the advantage that sample dilution is not
 237 required.

238 **Comparison of Transmission FT-NIR, Transmission FTIR,
 239 and SB-ATR/FTIR Methods for Wine Analysis.** A prelimi-
 240 nary comparative feasibility study was carried out by developing
 241 PLS calibrations for four parameters (alcohol content, total
 242 reducing sugars, total acidity, and pH) in wines by using
 243 transmission FTIR and FT-NIR as well as SB-ATR/FTIR spectra
 244 of 28 preanalyzed wine samples. The spectral regions employed
 245 in the optimized PLS calibration models for each of the four
 246 components are presented in **Table 1**. **Table 2** summarizes the
 247 leave-one-out cross-validation results in terms of the linear
 248 regression equations relating the cross-validation predictions to
 249 the values obtained by the reference methods, together with the
 250 corresponding values for R , SECV, MD_a , and SDD_a . The FT-
 251 NIR cross-validation data indicated poor agreement between
 252 the predicted and reference method values for all of the
 253 components except alcohol. Better overall agreement between
 254 the predicted and reference method values was obtained with
 255 both the transmission FTIR and SB-ATR/FTIR calibrations.
 256 Again, the best results were obtained for alcohol, whereas the

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257 results for reducing sugars, total acidity, and pH are very much
 258 in line with what has been reported in the literature for FTIR
 259 wine analysis using a transmission cell (16, 17).

260 The performance of the SB-ATR calibrations was generally
 261 comparable to that obtained with the conventional transmission
 262 cell, despite the much shorter effective path length of the SB-
 263 ATR accessory, and is in fact superior in the case of reducing
 264 sugars. This is likely due to the lower transmission cutoff of
 265 the ZnSe crystal as compared to CaF₂ windows, through which
 266 the transmission of IR energy declines rapidly between 1110
 267 and 1000 cm⁻¹, where the strongest absorption bands for sugars
 268 and alcohol are found (13). In addition, owing to the short
 269 effective path length of the SB-ATR accessory, all of the spectral
 270 information in the regions where water absorbs is available, and
 271 the CH stretching (3000–2800 cm⁻¹) and carbonyl stretching
 272 bands (1800–1600 cm⁻¹) are on scale. **Figure 2** illustrates the
 273 “pure component” spectra generated by the Turbo Quant Analyst
 274 software, each of which corresponds to the first loading spectrum
 275 in a PLS calibration for the component in question. The pure
 276 component spectra reveal that there is significantly less spectral
 277 information available in the transmission cell spectra. This is
 278 especially the case for total acidity, where the region between
 279 1750 and 1600 cm⁻¹ is totally “blacked out” due to off-scale
 280 water absorptions, whereas the SB-ATR spectra exhibit the
 281 characteristic carbonyl stretching absorption of carboxylic acids.

282 One consideration arising in the use of the ATR technique
 283 for quantitative analysis is the dependence of the effective path
 284 length on the depth of penetration (d_p) of the evanescent IR
 285 radiation emerging from the surface of the ATR crystal. The
 286 depth of penetration, in turn, is dependent on the refractive index
 287 of the sample in accordance with the equation

$$d_p = \lambda / \{2\pi n_1 [\sin^2(\theta) - (n_2/n_1)^2]^{1/2}\}$$

288 where λ is the wavelength of the IR radiation, n_1 is the refractive
 289 index of the ATR crystal, n_2 is the refractive index of the sample,
 290 and θ is the angle at which the IR radiation strikes the ATR
 291 crystal face (26). The effective path length thus may vary from
 292 sample to sample, depending on the refractive index of the
 293 sample material, whereas the calibration model is based on the
 294 assumption of a linear relationship between IR absorbance and
 295 concentration and, hence, a constant path length. In the case of
 296 distilled liquors, the value of the refractive index is almost
 297 invariant as the amount of alcohol in these types of products is
 298 almost always between 40 and 41% (v/v), with extremely low
 299 concentrations of other constituents. For wines, the refractive
 300 index is almost entirely dependent on the levels of alcohol and
 301 sugar, ranging from 1.34 for low-alcohol, dry wines to almost
 302 1.36 for high-sugar, high-alcohol wines. According to the
 303 equation above, for the SB-ATR accessory employed in the
 304 present study, which employs a ZnSe crystal and a 45° angle
 305 of incidence, this range corresponds to a variation of ~2.6% in
 306 the effective path length. Because the effective path length
 307 represented by the calibration model would correspond to an
 308 intermediate refractive index value, the errors in predicted
 309 concentrations resulting from this variability in the effective
 310 path length may be estimated to be <1.5%, which can be considered
 311 to be acceptable in relation to the errors presented in **Table 2**
 312 for the alternative FT-NIR and FTIR transmission methods.
 313 Overall, on the basis of the initial comparative results in **Table 2**,
 314 it was concluded that SB-ATR/FTIR spectroscopy has the
 315 potential to serve as a means by which multicomponent analysis
 316 of wines can be carried out.

317 **Multicomponent Wine Analysis by SB-ATR/FTIR Spec-**
 318 **troscopy.** Winemakers, enologists, and governments have an

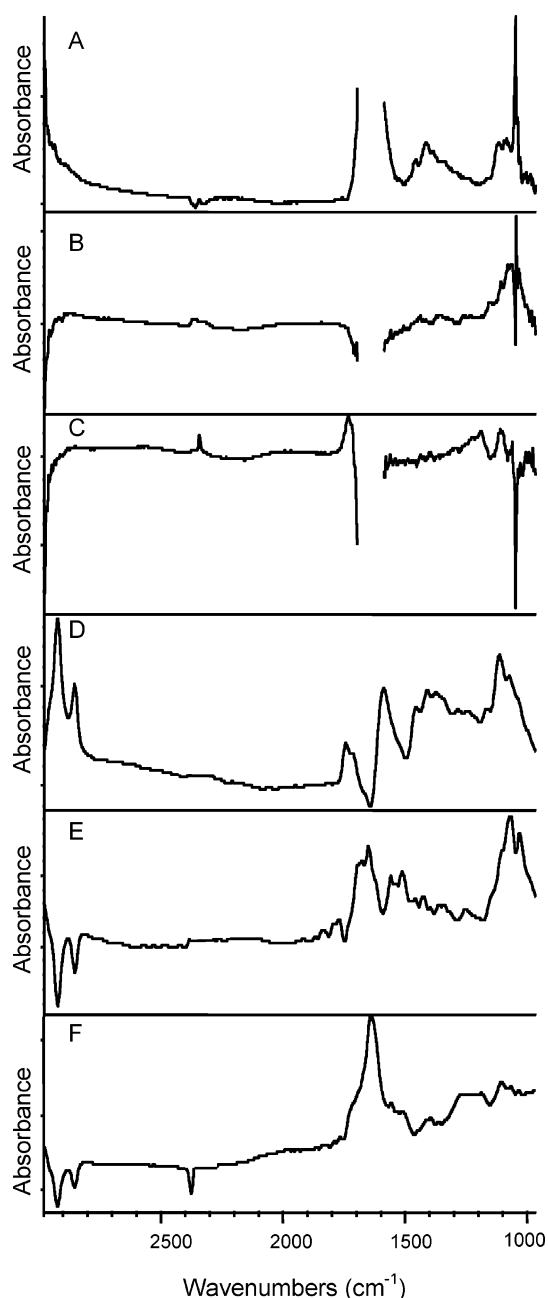


Figure 2. “Pure component” spectra derived for (A) pH, (B) total reducing sugar, and (C) total acidity from FTIR transmission spectra and for (D) pH, (E) total reducing sugar, and (F) total acidity from SB-ATR spectra of 28 wine samples.

319 interest in a variety of components and parameters that affect
 320 the quality of wine, are indicative of adulteration or provenance,
 321 or affect revenue. In this study, PLS calibrations were developed
 322 for a total of 11 components and parameters using a diverse set
 323 of 149 wine samples, for which reference values were obtained
 324 by combining data from the Société des alcools du Québec and
 325 HPLC analyses performed in our laboratory. About half of the
 326 samples were employed as calibration standards, with the
 327 balance serving as an independent validation set.

328 In the development of the PLS calibration models, five broad
 329 spectral regions were initially selected by examining the pure
 330 component spectra. The optimal spectral region or combination
 331 of spectral regions for the prediction of each of the 11 parameters
 332 was chosen by evaluating the cross-validation statistics. The
 333 regions ultimately selected are summarized in **Table 3**. The
 334 optimized calibration models were validated by comparing the

Table 3. Regions Employed in PLS Regressions for the Analysis of 11 Components in 72 Wines by SB-ATR/FTIR Spectroscopy

component	region				
	1500–850 cm ⁻¹	1500–957 cm ⁻¹	1500–1108 cm ⁻¹	1801–1500 cm ⁻¹	3000–2800 cm ⁻¹
alcohol	+	–	–	–	–
fructose	–	+	–	–	+
citric acid	+	–	–	+	+
tartaric acid	+	–	–	+	+
malic acid	–	–	+	+	–
glycerol	+	–	–	+	–
lactic acid	+	–	–	+	–
reducing sugar	–	+	–	–	+
volatile acidity	–	+	–	+	–
total acidity	–	–	+	+	–
total SO ₂	+	–	–	–	–

335 PLS-predicted values for the 77 preanalyzed wines in the
 336 validation set to the reference method values. **Table 4** sum-
 337 marizes the data obtained for all 11 parameters, including the
 338 ranges of the values spanned by the validation set and the linear
 339 regression equations relating the PLS predicted values to the
 340 reference method values as well as the values for *R*, SEP, MD_a,
 341 and SDD_a. For purposes of comparison, the SEP and SDD_a
 342 values reported by Patz et al. (16) and Dubernet and Dubernet
 343 (17), respectively, for FTIR wine analysis using a commercial
 344 wine analyzer are also included in **Table 4**.

345 The predicted values for the parameters evaluated in the initial
 346 trial (alcohol, reducing sugars, and total acidity) corresponded
 347 well to the reference method values for the validation samples;
 348 compared to the initial trial set in **Table 2**, higher *R* values
 349 were obtained (>0.90) because the data set in **Table 4** spanned
 350 a larger range of concentration values. A high correlation value
 351 (0.987) was also obtained for fructose. It may be noted that the
 352 validation set was uniformly distributed over the entire range
 353 of alcohol content, whereas for sugars, most of the wines were
 354 dry with only a few of them being semisweet or sweet. The
 355 possibility of simplifying the calibration process by developing
 356 PLS calibration models for the prediction of these two param-
 357 eters based simply on mixtures of ethanol, glycerol, fructose,
 358 and glucose as calibration standards was investigated. However,
 359 large biases were obtained when wines were analyzed using
 360 these calibrations, indicating that these simple calibration
 361 standards did not adequately model the spectra of wines. This
 362 finding is consistent with results obtained by Schindler et al.
 363 (4) in their investigation of a calibration set consisting of model
 364 solutions as an alternative to preanalyzed wines.

365 Satisfactory agreement between the FTIR-predicted and
 366 reference values was also obtained for glycerol, a component

present in wines as a fermentation byproduct (27), and for the
 individual organic acids, with the exception of citric acid. Better
 results were obtained for total acidity (*R* = 0.925) than for the
 individual organic acids, which may be attributed to differences
 in the precision of the reference methods as well as to possible
 inadequacy of the FTIR method in differentiating among the
 various organic acids. The poor results for citric acid are due
 to its very low concentration, as naturally occurring levels in
 wine are typically below 0.1 g L⁻¹ (27), and very few of the
 samples available for this study contained >0.30 g L⁻¹.
 However, citric acid is sometimes added to wines to increase
 the total acidity content or to prevent casse, a winemaking term
 used to describe cloudiness or formation of a precipitate due to
 an excess of certain metal salts (27). If quantitation of citric
 acid in these types of samples is required, a calibration model
 based on wines spiked with known amounts of citric acid to
 extend the concentration range and enhance the spectral response
 can be developed.

In addition to citric acid, volatile acidity and total SO₂ were
 poorly predicted. Volatile acidity levels were too low and
 spanned too narrow a range of concentration (0.2–0.8 g L⁻¹
 H₂SO₄) to develop an adequate calibration, whereas total SO₂,
 present at the parts per million level in wines, does not produce
 an adequate signal to calibrate on at concentrations of <100
 ppm (16), and its spectrum is complicated by its multiple ionic
 states, which are sensitive to the sample matrix pH (17).

A detailed comparison between the SEP and SDD_a values
 obtained in this study and the values from the literature given
 in **Table 4** would not be informative because the analytical
 performance achieved for each component is highly dependent
 on the precision of the reference method used to analyze the
 calibration and validation samples and the number and types
 of wine samples used in the calibration and validation sets.
 However, overall, the data in **Table 4** indicate that the SB-
 ATR/FTIR results are comparable to those obtained in studies
 using a commercial FTIR wine analyzer equipped with a CaF₂
 flow-through cell (16, 17) (**Table 4**). Given the results in **Table**
2 for the SB-ATR accessory and a 50-μm CaF₂ transmission
 cell, which can be directly compared with each other because
 they are based on a common sample set, and in **Table 4**, it
 may be concluded that SB-ATR/FTIR spectroscopy can provide
 comparable sensitivity to commercial FTIR wine analyzers,
 capable of measuring concentrations of components at levels
 of >0.1 g L⁻¹ (16, 17).

Conclusion. This study has demonstrated the suitability of
 SB-ATR/FTIR spectroscopy for the analysis of distilled liquors
 and wines. The determination of alcohol in distilled liquors is
 a simple analytical procedure, and calibration can be performed

Table 4. Linear Regression Equations Relating PLS Predictions to the Reference Values for a Validation Set of 77 Wine Samples Ranked in Terms of Decreasing *R* Value with Comparison of SEP and SDD_a to Data Published from Studies Using a CaF₂ Transmission Flow-through Cell

component	range of concentration	equation	<i>R</i>	SEP	SEP ^a	MD _a	SDD _a	SDD _a ^b
alcohol	7.3–17.0% (v/v)	Y = -0.235 + 1.020X	0.997	0.099	0.21	0.01	0.101	0.08
reducing sugar	1.3–116.6 g L ⁻¹	Y = -0.729 + 0.970X	0.992	2.180	0.86	-0.94	2.231	2.41
fructose	0–64.48 g L ⁻¹	Y = -0.179 + 1.040X	0.987	1.606	0.62	-0.08	1.630	
total acidity	2.7–6.3 g L ⁻¹ (H ₂ SO ₄)	Y = 0.172 + 0.906X	0.925	0.178	0.13 ^c	-0.17	0.182	0.07
lactic acid	0.03–3.47 g L ⁻¹	Y = 0.366 + 0.821X	0.901	0.291	0.20	0.12	0.317	0.30
glycerol	2.07–15.53 g L ⁻¹	Y = 2.136 + 0.796X	0.864	0.824	0.49	0.84	0.894	1.315
malic acid	0–4.54 g L ⁻¹	Y = 0.498 + 0.700X	0.768	0.651	0.16	0.26	0.727	0.12
tartaric acid	0.43–3.81 g L ⁻¹	Y = 0.432 + 0.764X	0.709	0.388	0.38	-0.09	0.404	0.11
volatile acidity	0.21–0.83 g L ⁻¹ (H ₂ SO ₄)	Y = 0.281 + 0.472X	0.625	0.105	0.071 ^c	0.04	0.14	0.022
total SO ₂	0–250 ppm	Y = 54.27 + 0.307X	0.569	21.1	17	-5.9	38.8	
citric acid	0–3.37 g L ⁻¹	Y = 0.126 + 0.081X	0.384	0.104		-0.11	0.497	

^a SEP values from Patz et al. (16). ^b SDD_a values from Dubernet and Dubernet (17). ^c SEP values for total acidity and volatile acidity in Patz et al. (16) expressed as g L⁻¹ tartaric acid and g L⁻¹ acetic acid, respectively.

415 using a small set of ethanol/distilled water mixtures as standards.
 416 In contrast, multicomponent wine analysis is a complex
 417 undertaking, requiring a very careful selection of the wines used
 418 as calibration standards to represent the population of interest
 419 and substantial investment in quality reference method analyses.
 420 This drawback pertains equally to commercially available FTIR
 421 wine analyzers (16, 17). In this context, the use of gravimetri-
 422 cally prepared solutions containing the pure components of
 423 interest as calibration standards would clearly be advantageous.
 424 However, an investigation of this possibility in the present work
 425 indicated that such an approach was not viable owing to the
 426 complexity of the wine matrix, corroborating the findings of a
 427 previous study (4).

428 As an alternative to the transmission cell employed in FTIR
 429 wine analyzers, the SB-ATR accessory yielded comparable
 430 analytical performance. In addition, it provided a very simple
 431 means of sample handling, as only a drop of sample is required
 432 to cover the surface of the ATR crystal for analysis and the
 433 sample can simply be wiped off the crystal, which is then ready
 434 for the next analysis after a quick rinse with water. Furthermore,
 435 the SB-ATR sample-handling accessory can be adapted for
 436 automated flow-through analysis, which would avoid the need
 437 to cover the crystal to prevent evaporation and to clean the
 438 surface of the crystal between samples. Generally speaking, this
 439 method provides a means of analyzing a wide range of alcoholic
 440 beverages, including products high in alcohol and/or sugar
 441 content without any need for prior dilution of the sample. The
 442 proven suitability of an SB-ATR accessory for wine analysis
 443 makes it possible to readily implement FTIR wine analysis
 444 methodology on a standard benchtop FTIR spectrometer,
 445 furnishing a less costly alternative to the purchase of a dedicated
 446 analyzer. Furthermore, the standard SB-ATR accessory can be
 447 adapted to a flow-through configuration to accommodate high-
 448 volume analytical applications in distilleries, wineries, and
 449 commercial and regulatory laboratories.

450 ABBREVIATIONS USED

451 ATR, attenuated total reflectance; FTIR, Fourier transform
 452 infrared; FT-NIR, Fourier transform near-infrared; HPLC, high-
 453 performance liquid chromatography; IR, infrared; MD_a, mean
 454 difference for accuracy; NIR, near-infrared; PLS, partial least
 455 squares; *R*, coefficient of correlation; SB-ATR, single-bounce
 456 attenuated total reflectance; SDD_a, standard deviation of the
 457 differences for accuracy; SECV, standard error of cross-
 458 validation; SEP, standard error of prediction.

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463 LITERATURE CITED

- 464 (1) *Encyclopedia of Food Science, Food Technology and Nutrition*;
 465 Macrae, R., Ed.; Academic Press: San Diego, CA, 1993; pp
 466 4921–4941.
 467 (2) Amerine, M. A.; Ough, C. S. In *Methods for Analysis of Musts*
 468 *and Wines*, 1st ed.; Wiley: New York, 1980.
 469 (3) Davenel, A.; Grenier, P.; Foch, B.; Bouvier, J. C.; Verlaque, P.;
 470 Pourcin, J. Filter, Fourier transform infrared, and areometry, for
 471 following alcoholic fermentation in wines. *J. Food Sci.* **1991**,
 472 *56* (6), 1635–1638.
 473 (4) Schindler, R.; Vonach, R.; Lendl, B.; Kellner, R. A rapid
 474 automated method for wine analysis based upon sequential
 475 injection (SI)-FTIR spectrometry. *Fresenius' J. Anal. Chem.*
 476 **1998**, *362*, 130–136.

- (5) Li-Chan, E. C. Y.; Ismail, A. A.; Sedman, J.; van de Voort, F. 477
 R. Vibrational spectroscopy of food and food products. In 478
Handbook of Vibrational Spectroscopy; Chalmers, J. M., Grif- 479
 fiths, P. R., Eds.; Wiley: Chichester, U.K., 2002; pp 3629– 480
 3662. 481
 (6) Cabanis, M. T.; Cabanis, J. C.; Raffy, J.; Peyronnenche, M.; 482
 Viotte, O.; Leboeuf, J. P. L'Infralyzer 400: une détermination 483
 automatique du titre alcoolométrique des vins. *Rev. Fr. Oenol.* 484
1983, *89*, 75–79. 485
 (7) van den Berg, F. W. J.; van Osenbruggen, W. A.; Smilde, A. K. 486
 Process analytical chemistry in the distillation industry using 487
 near-infrared spectroscopy. *Process Control Qual.* **1997**, *9*, 51– 488
 57. 489
 (8) Gallignani, M.; Garrigues, S.; de la Guardia, M. Direct deter- 490
 mination of ethanol in all types of alcoholic beverages by near- 491
 infrared derivative spectrometry. *Analyst* **1993**, *118*, 1167–1173. 492
 (9) Tipparat, P.; Lapanantnoppakhum, S.; Jakmunee, J.; Grudpan, 493
 K. Determination of ethanol in liquor by near-infrared spectro- 494
 photometry with flow injection. *Talanta* **2001**, *53*, 1199–1204. 495
 (10) Garcia-Jares, C. M.; Medina, B. Application of multivariate 496
 calibration to the simultaneous routine determination of ethanol, 497
 glycerol, fructose, glucose and total residual sugars in botrytized- 498
 grape sweet wines by means of near-infrared reflectance 499
 spectroscopy. *Fresenius' J. Anal. Chem.* **1997**, *357*, 86–91. 500
 (11) Damberg, R. G.; Kambouris, A.; Francis, I. L.; Gishen, M. Rapid 501
 analysis of methanol in grape-derived distillation products using 502
 near-infrared transmission spectroscopy. *J. Agric. Food Chem.* 503
2002, *50*, 3079–3084. 504
 (12) Urbano-Cuadrado, M.; Luque de Castro, M. D.; Pérez-Juan, P. 505
 M.; Garcia-Olmo, J.; Gomez-Nieto, M. A. Near infrared reflectance 506
 spectroscopy and multivariate analysis in enology. Deter- 507
 mination or screening of fifteen parameters in different types of 508
 wines. *Anal. Chim. Acta* **2004**, *527*, 81–88. 509
 (13) Bouvier, J. C. Réflexion sur l'analyse oenologique par spec- 510
 trométrie infrarouge. *Rev. Fr. Oenol.* **2001**, *191*, 16–17. 511
 (14) Mahia, P. L.; Gandara, J. S.; Losada, P. P. Validation of ethanol 512
 determination in alcoholic beverages by infrared spectro- 513
 photometry using orthogonal and derivative functions to correct 514
 for water absorption. *Vibr. Spectrosc.* **1992**, *3*, 133–138. 515
 (15) Gallignani, M.; Garrigues, S.; de la Guardia, M. Derivative 516
 Fourier transform infrared spectrometric determination of ethanol 517
 in alcoholic beverages. *Anal. Chim. Acta* **1994**, *287*, 275–283. 518
 (16) Patz, C. D.; David, A.; Thente, K.; Kurbel, P.; Dietrich, H. Wine 519
 analysis with FTIR spectrometry. *Vitic. Enol. Sci.* **1999**, *54* (2– 520
 3), 80–87. 521
 (17) Dubernet, M.; Dubernet, M. Utilisation de l'analyse infrarouge 522
 à transformée de Fourier pour l'analyse oenologique de routine. 523
Rev. Fr. Oenol. **2000**, *181*, 10–13. 524
 (18) Andersen, S. T.; Hansen, P. W.; Andersen, H. V. Vibrational 525
 spectroscopy in the analysis of dairy products and wine. In 526
Handbook of Vibrational Spectroscopy; Chalmers, J. M., Griffiths, 527
 P. R., Eds.; Wiley: Chichester, U.K., 2002; pp 1–10. 528
 (19) Brierley, P. R. Imaging stage for Fourier transform infrared 529
 spectrometer. U.S. Patent 5,965,889, 1999. 530
 (20) Cocciardi, R. A.; van de Voort, F. R.; Sedman, J.; Ismail, A. A. 531
 Monitoring of lactose hydrolysis in milk by single-bounce 532
 attenuated total reflectance Fourier transform infrared spectro- 533
 scopy. *Milchwissenschaft* **2004**, *59* (7–8), 403–407. 534
 (21) Association of Official Analytical Chemists. In *Official Methods* 535
of Analysis, 16th ed.; AOAC International: Arlington, VA, 1995; 536
 sec. 982.10. 537
 (22) Haaland, D. M.; Thomas, E. V. Partial-least-squares methods 538
 for spectral analyses. I. Relation to other quantitative calibration 539
 methods and the extraction of qualitative information. *Anal.* 540
Chem. **1988**, *60*, 1193–1201. 541
 (23) Association of Official Analytical Chemists. In *Official Methods* 542
of Analysis, 16th ed.; AOAC International: Arlington, VA, 1995; 543
 sec. 942.06. 544
 (24) Association of Official Analytical Chemists. In *Official Methods* 545
of Analysis, 16th ed.; AOAC International: Arlington, VA, 1995; 546
 sec. 950.04. 547

548	(25) Association of Official Analytical Chemists. In <i>Official Methods of Analysis</i> , 16th ed.; AOAC International: Arlington, VA, 1995; sec. 957.03.	In <i>The Technology of Wine Making</i> , 4th ed.; Avi Publishing: Westport, CT, 1980; pp 715–755.	557 558
550	(26) Noiseux, I.; Long, W.; Courmoyer, A.; Vernon, M. Simple fiber-optic-based sensors for process monitoring: an application in wine quality control monitoring. <i>Appl. Spectrosc.</i> 2004 , 58 (8), 1010–1019.		559
551	(27) Amerine, M. A.; Berg, H. W.; Kunkee, R. E.; Ough, C. S.; Singleton, V. L.; Webb, A. D. Legal restrictions on wine making.	Received for review August 6, 2004. Revised manuscript received January 5, 2005. Accepted January 12, 2005. Financial support for this work by Thermal-Lube Inc. is gratefully acknowledged.	560 561 562
552		JF048663D	563