

# Chemometric Profiling of Whiskey Using the 5977A GC/MSD

### **Application Note**

Food Testing & Agriculture

### Abstract

Nontargeted compound analysis and statistical tools were used in combination with the high sensitivity of the Agilent 5977A Series GC/MSD with Extractor El Source to generate compound profiles that were used to differentiate five brands of whiskey.

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### Introduction

Gas chromatography/mass spectrometry (GC/MS) is widely used in food analysis for applications such as R&D, quality control, and quality assurance. Advances in GC/MS performance have enabled reliable detection of the myriad of trace compounds common to most natural products. Although human sensory tests (smell and taste) are still an essential part of flavor quality control, GC/MS is able to provide increasingly valuable details about changes and differences in the concentration profiles of major and trace components without the limitation of human sensors.

Chemometrics can be used to solve both descriptive and predictive problems. In descriptive applications, properties of chemical systems are modeled with the intent of learning the underlying relationships and structure of the system. In predictive applications, properties of chemical systems are modeled to predict new properties or behavior of interest. GC/MS is often used to derive the data used in both descriptive and predictive chemometrics. In the predictive mode, this technique has been used to predict whether olive oil will pass the extra virgin sensory test [1], distinguishing wine [2] varieties, and whether shochu is contaminated during the manufacturing process [3]. In the descriptive mode, this technique can be used to distinguish closely related food products, such as different brands of whiskey.

While these chemometric analyses are often performed using very powerful MS instrumentation, lower cost single quadrupole mass detectors can also provide useful information. This application note demonstrates the use of sophisticated statistical analysis of data generated by the 5977A GC/MSD to distinguish differences between five different brands of whiskey. The 5977A GC/MSD, in combination with the Agilent 7890B GC, is an ideal platform for sensitive and sophisticated statistical profiling of food products such as whiskeys, and automated solid phase micro-extraction (SPME) on the PAL Automated Sample Injector enables very sensitive headspace sampling of the whiskey aromas. Mass Profiler Professional (MPP) Software enables classification of the composition of complex samples such as whiskey using a range of statistical tools.

This study used nontargeted compound analysis and statistical tools such as one-way analysis of variance (ANOVA), principal component analysis (PCA) and hierarchical cluster analysis (HCA) to identify differences between the various brands of whiskeys. Data and statistical analyses were performed using NIST AMDIS (Automated Mass Spectral Deconvolution and Identification System), Agilent MassHunter ID Browser and Mass Profiler Professional software. This approach enabled classification of the whiskeys into four groups based on the relative concentrations of 46 different entities.

### **Experimental**

### Samples

Five different whiskeys were obtained commercially in the US, and they are described in Table 1.

| Sample             | Description   | Subjective aroma   |  |
|--------------------|---|--|--|
| Popular brand (PB) | Most popular whiskey in the market  | A soft, thin entry to an off-dry   |  |
| Competitor (A)     | Described as premium whiskey  | Similar to PB  |  |
| Competitor (B)     | Popular knock-off whiskey   | Sweet with light caramel and vanilla flavors. Stronger aroma than PB   |  |
| Competitor (C)     | Claims to be even higher quality than PB Sweet aroma, slightly stronger than PB |  |  |
| Competitor (D)     | Claims to be a deep flavor whiskey  | ims to be a deep flavor whiskey Honey, butter, and a hint of dark fruit (plums, raisins). Stronger aroma than PB |  |

Table 1. Whiskey Samples Used in the Study

### Instruments

This study was performed on an Agilent 7890B GC equipped with automated solid phase micro-extraction (SPME) on the PAL Automated Sample Injector and coupled to the single quadrupole Agilent 5977A GC/MSD with Extractor El Source. The instrument conditions are listed in Tables 2 and 3.

### **Sample preparation**

The volatile odor and flavor components from each sample type were collected using headspace SPME. Each 5 mL whiskey sample was transferred to a 10 mL headspace vial. A 50  $\mu$ m × 2 cm DVB/CAR/PDMS was exposed to the headspace of the sample at 60 °C for 10 minutes with agitation. Volatile compounds absorbed on the SPME fiber were thermally desorbed at 240 °C for 1 minute into an injection port.

Table 2. PAL Automated Sample Injector SPME Conditions

| Sample volume                 | 5 mL of whiskey in a 10 mL vial       |  |  |
|-------------------------------|---------------------------------------|--|--|
| Syringe                       | 2 cm Fiber 50/30 $\mu m$ DVB/CAR/PDMS |  |  |
| Pre-incubation time           | 60 seconds                            |  |  |
| Incubation temperature        | 60 °C                                 |  |  |
| Pre-incubation agitator speed | 500 rpm                               |  |  |
| Agitator time                 | On at 0 seconds, off at 2 seconds     |  |  |
| Vial needle penetration       | 11 mm                                 |  |  |
| Vial fiber exposure           | 22 mm                                 |  |  |
| Extraction time               | 600 seconds                           |  |  |
| Desorb to                     | Split/splitless inlet                 |  |  |
| Injection needle penetration  | 32 mm                                 |  |  |
| Injection fiber exposure      | 22 mm                                 |  |  |
| Desorption time               | 60 seconds                            |  |  |
|                               |                                       |  |  |

Table 3. GC and Mass Spectrometer Conditions

### GC run conditions

| GC run conditions                               |   |  |  |
|---|---|--|--|
| Analytical column                               | HP INNOWAX (25 m × 0.20 mm, 0.40 µm) (p/n 19091N-202)                                     |  |  |
| Injection method                                | SPME (50/30 µm DVB/CAR/PDMS)  |  |  |
| Inlet temperature                               | emperature Isothermal at 260 °C   |  |  |
| Injection mode                                  | Split, 50:1 ratio   |  |  |
| Oven temperatures                               | 1.5 minutes hold at 40 °C<br>40 °C to 240 °C at 30 °C/min<br>Hold at 240 °C for 3 minutes |  |  |
| Column flow                                     | 1.1 mL/min constant flow  |  |  |
| Carrier gas Helium                              |   |  |  |
| Transfer line temp                              | 255 °C  |  |  |
| GC run time                                     | 16 minutes  |  |  |
| MS conditions                                   |   |  |  |
| Ionization mode                                 | EI, 70 eV   |  |  |
| lon source temperature                          | 230 °C  |  |  |
| Quadrupole temperature 150 °C                   |   |  |  |
| Acquisition mode Scan (50–550 amu), normal mode |   |  |  |
| A/D sample                                      | 4   |  |  |
| EM setting gain                                 | 1.0   |  |  |
| Threshold                                       | 150   |  |  |
| Trace ion detection On                          |   |  |  |
| Tuning  | etune.u and atune.u   |  |  |

### Data processing and statistical analysis

Entity extraction from the GC/MS data was done using AMDIS on the Agilent MSD Productivity ChemStation (F.01.00). The .ELU files from AMDIS were imported into Mass Profiler Professional (MPP) for differential analysis. MPP 12.1 was used for data filtering and statistical analysis, and compound identification was performed using the NIST 11 MS Library and Agilent MassHunter ID Browser. The settings used for these software packages are shown in Table 4.

### **Results and Discussion**

## Detection of trace compounds, Etune versus Atune

The 5977A GC/MSD features a unique Extractor EI Source and its Etune tuning protocol, which increase MSD sensitivity in order to achieve lower detection limits and improve the identification of trace-level compounds. The Atune algorithm from previous generations of the Agilent MSD is still available for use with the Extractor EI Source. Both tuning protocols were used in the detection of trace compounds from the aromas of the whiskey samples, in order to compare their relative efficiencies for this application. Table 4. Data Processing and Statistical Analysis Software Settings

#### **Deconvolution (AMDIS 2.67)**

| 12   |  |  |  |  |
|--|--|--|--|--|
| 0 (TIC), 207, 267  |  |  |  |  |
| Тwo  |  |  |  |  |
| Medium   |  |  |  |  |
| Low  |  |  |  |  |
| Medium   |  |  |  |  |
| er Professional 12.1)  |  |  |  |  |
| > 20   |  |  |  |  |
| > 1,000  |  |  |  |  |
| > 3  |  |  |  |  |
| < 0.10   |  |  |  |  |
| > 0.3  |  |  |  |  |
| None   |  |  |  |  |
| Compound identification<br>(NIST MS Library and Agilent MassHunter ID Browser) |  |  |  |  |
| NIST 11  |  |  |  |  |
| > 50, Best hit   |  |  |  |  |
|  |  |  |  |  |

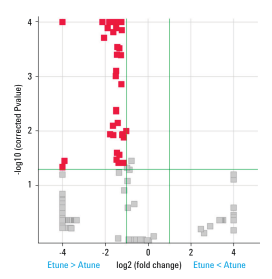
Analyzing samples using Etune and Atune generated a combined list of 142 entities from four replicate injections. A comparison of the relative intensities revealed that 48 of the 142 entities with a fold change  $\geq 2$  between the two tuning protocols passed the t-test at a probability p-value < 5%, as shown in red in the volcano plot in Figure 1. All 48 exhibited higher intensities with Etune, versus Atune. In fact, four entities found using Etune were not detected using Atune, under the same AMDIS parameters (Figure 2).

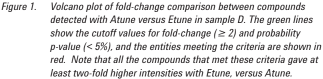
### Profiling of whiskey aroma compounds

In order to fully characterize the compounds constituting the aroma of the five whiskey samples, GC/MS analysis was conducted in triplicate on all five whiskey samples. The detected entities were then filtered using a coefficient of variation (CV) filter of noise reduction of 75%, resulting in 74 entities common to the five whiskey samples. These were then divided into two groups (Figure 3), those with relative peak intensities <1,000,000 (low and medium abundance), and those with peak intensities  $\geq$  1,000,000 (high abundance).

| Retention time | Mass  | Compound                     | Atune                                  | Etune                            |
|----------------|-------|------------------------------|--|----------------------------------|
| 6.522          | 401   | 401.0 at 6.52155             | 0                                      | 27,685                           |
| 11.415         | 88    | Pentadecanoic acid, ethyl es | ter O                                  | 25,542                           |
| 7.86           | 475   | 475.0 at 7.86025             | 0                                      | 18,527                           |
| 8.646          | 143   | 143.0 at 8.646299            | 0                                      | 19,100                           |
| 6,500          |       |                              | 00 (97 70 -                            | 88.70): D-et                     |
| 6,000          |       |                              |  | 88.70): D-et                     |
|                |       | EIC 88                       | .00 (87.70 ~                           | · 88.70): D-et                   |
| 5,500          |       |                              |  | 88.70): D-et                     |
| 5,000          |       |                              |  | · 88.70): D-at<br>· 88.70): D-at |
|                |       | EIC 88                       | .00 (87.70 ~                           | 88.70): D-at                     |
| 1,500          |       | Etune EIC 88                 | .00 (87.70 ~                           | 88.70): D-at                     |
| 1,000          |       | h                            |  |                                  |
| 3,500          |       | A                            |  |                                  |
|                |       |                              |  |                                  |
| 3,000          |       | //                           |  |                                  |
| 2,500          |       |                              |  |                                  |
|                |       |                              |  | Ann                              |
| 2,000          | ۵     |                              | Marc                                   | [ And ]                          |
| 1,500          | Λ ~   |                              | V XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX | $\varphi \circ v_{\bullet}$      |
| 1,000          | / 165 |                              |  |                                  |
| -              | A -   | A                            |  |                                  |
| 500            |       | Atuno                        | •                                      |                                  |
| 04             |       | Atune                        |  |                                  |
| 11.00 11.      | 10 11 | .20 11.30 11.40 11.50        | 11.60 11                               | .70 11.80                        |
|                |       | Time                         |  |                                  |

Figure 2. Four compounds were detected in four replicate analyses of sample D using Etune (upper chart) that were not seen when using Atune and the same AMDIS integration threshold. The lower extracted ion chromatograms (EIC) show the 88u peak (pentadecanoic acid, ethyl ester) in the four replicates, using Etune and Atune.





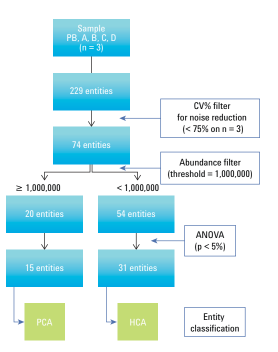


Figure 3. The workflow for chemometric profiling of the whiskey samples, culminating in classification of the relevant compounds by principal component analysis (PCA) or hierarchical cluster analysis (HCA).

### **High abundance entities**

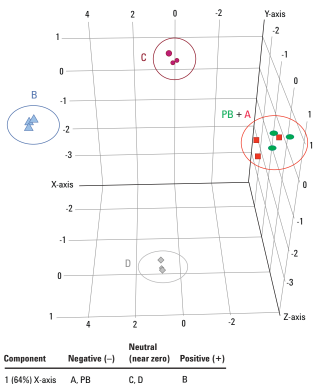
Twenty compounds with peak intensities  $\geq$  1,000,000 were identified using Mass Profiler Professional (MPP). For comparisons of the differences in the tuning, both Atune and Etune were again used. Filtering these using one-way

analysis of variance (ANOVA) with a p-value < 5% resulted in 15 compounds of statistical significance (Figure 4). Principal component analysis (PCA) of these 15 resulted in four groups of distinguishable samples, PB+A, B, C, and D (Figure 5).

| Retention Time | Mass  | Compound                           | [PB]    | [A]      | [B]      | [C]      | [D]      |
|----------------|-------|------------------------------------|---------|----------|----------|----------|----------|
| 2.590          | 61.0  | Trimethylsilylmethanol             | 3759228 | 4107974  | 3257738  | 6529116  | 8362442  |
| 4.515          | 74.0  | 2-Amino-2-methyl-1,3-propanediol   | 3863118 | 3302027  | 1622565  | 2550368  | 1281017  |
| 4.772          | 70.0  | 1-Butanol, 3-methyl-, acetate      | 3286931 | 3442918  | 1255734  | 584678   | 384032   |
| 5.724          | 88.0  | Hexanoic acid, ethyl ester         | 855511  | 840595   | 2161864  | 991785   | 1130796  |
| 7.246          | 88.0  | Octanoic acid, ethyl ester         | 6435439 | 7319735  | 35215980 | 7083341  | 16339178 |
| 7.590          | 96.0  | 3-Furaldehyde                      | 3406372 | 3151482  | 1785684  | 1587822  | 3265564  |
| 7.937          | 88.0  | Nonanoic acid, ethyl ester         | 619252  | 708628   | 1281332  | 674696   | 716060   |
| 8.593          | 88.0  | Decanoic acid, ethyl ester         | 9160368 | 10945965 | 91484704 | 15420839 | 43557596 |
| 8.721          | 70.0  | Octanoic acid, 3-methylbutyl ester | 99822   | 132939   | 1301459  | 366510   | 160848   |
| 8.767          | 110.0 | Ethyl trans-4-decenoate            | 528019  | 555572   | 1170285  | 717243   | 240995   |
| 9.803          | 88.0  | Dodecanoic acid, ethyl ester       | 1428005 | 1770075  | 31862326 | 3835626  | 14279053 |
| 9.912          | 70.0  | (-)-1-Methylbutyl decanoate        | 35299   | 45598    | 1048773  | 303444   | 187453   |
| 10.348         | 91.0  | Phenylethyl Alcohol                | 1451716 | 1323375  | 2749470  | 2825832  | 1035489  |
| 10.901         | 88.0  | Tetradecanoic acid, ethyl ester    | 183026  | 188006   | 3107707  | 483733   | 1213085  |
| 11.906         | 88.0  | Hexadecanoic acid, ethyl ester     | 343726  | 300003   | 2559052  | 741498   | 1686823  |

Figure 4. Fifteen high abundance compounds identified using MPP and filtered using ANOVA with a p-value < 5% to assure statistical significance.

In Component 1, sample B has a high positive score in the PCA Score Plot and sample A and PB have negative scores (Figure 5). The entities are located in the PCA Loading Plot (Figure 6) according to the loading of Components 1 and 2. In the PCA Loading Plot, the entities that are unique to sample B are placed on the positive loading of Component 1. By comparing the PCA Score Plot (Figure 6A) and the Loading Plot (Figure 6B), we can identify unique entities that differentiate the various whiskey samples. The table in Figure 6 shows the example of the entities that are unique to sample B, and these are the entities that provide sample B with a high score for Component 1, which is the x-axis in the PCA Loading Plot (Figure 5). Also, the relative peak intensities of the components of each group are characteristic of that group (Figure 6).



| 1 (64%) X-axis | A, PB | C, D      | В           |
|----------------|-------|-----------|-------------|
| 2 (22%) Y-axis | D     | _         | A, B, C, PB |
| 3 (11%) Z-axis | С     | -         | A, B, D, PB |
|                |       | C .1 1E . | · · · · · · |

Figure 5. PCA analysis of the 15 significantly relevant compounds in the high abundance group results in four distinctive groups of compounds that differentiate all of the samples except A and B.

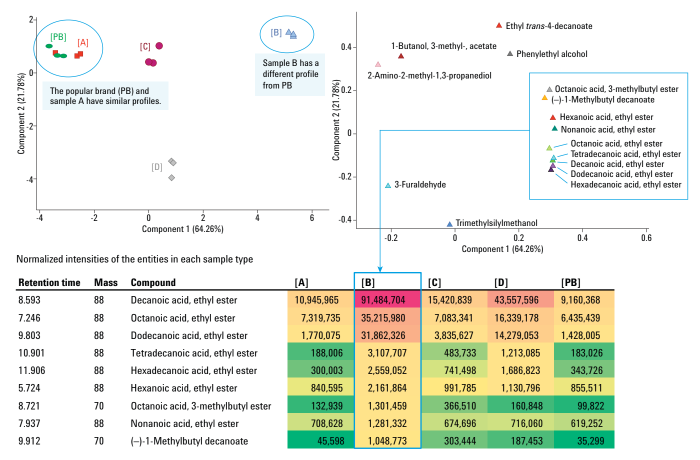


Figure 6. PCA scores illustrate the separation of the four sample groups (upper score plots), and the relative normalized intensities of the components of each group are characteristic for that group (lower table). Some of the components in the PCA Loading Plot are overlapped because they have similar profiles. Red: very high intensity; Orange: high intensity; Yellow: moderate intensity; Green: low intensity.

### Low and medium abundance entities

Fifty-four entities with peak intensities <1,000,000 were detected using Etune and identified using Mass Profiler Professional (MPP). Filtering these using ANOVA with a p-value <5% resulted in 31 compounds of statistical significance. Hierarchical Cluster Analysis (HCA) of these 31 again grouped the samples into four groups, PB+A, B, C, and D (Figure 7). In turn, the 31 compounds were classified by HCA according to the similarity of the normalized intensity profiles into eight clusters, which are shown in detail in Figure 8.

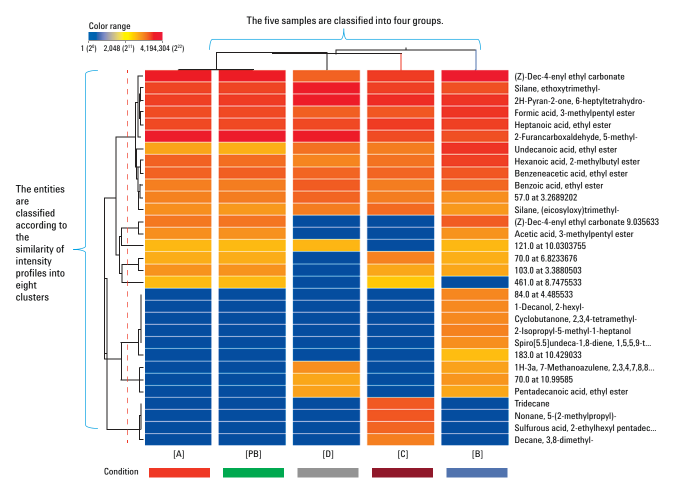
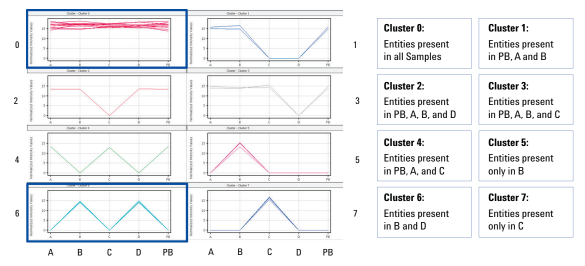
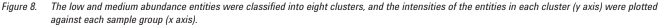


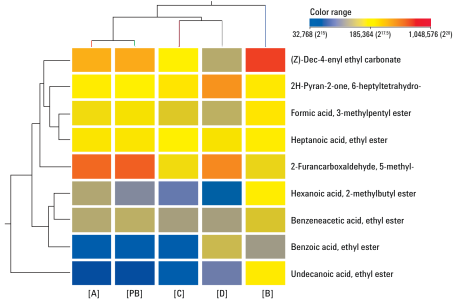
Figure 7. Classification of the samples and entities using those entities with normalized intensities < 1,000,000 and HCA. The samples were classified into four groups, and the entities were grouped into eight clusters according to profile similarity.

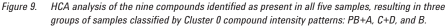
Of the 12 entities present in all five samples (Cluster 0), one was unidentified and another two were system blanks. The remaining nine were classified into three groups of distinguishable samples by entity intensity: PB+A, C+D, and B (Figure 9). The normalized intensities ranged from 14,000 to 500,000.

Samples B and D both contained three entities in Cluster 6 (Figure 10). One of these was pentadecanoic acid ethyl ester, which had been identified earlier in Sample D in the comparison of the Atune and Etune protocols. This compound was only found using Etune.









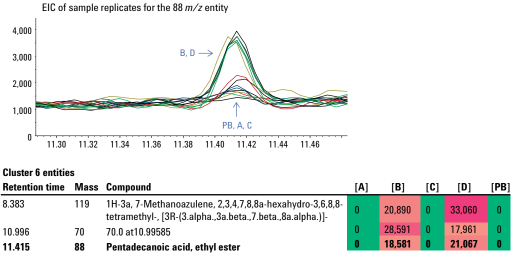


Figure 10. Identities and normalized intensities of the Cluster 6 entities, as well as extracted ion chromatograms (EICs) for the replicate analyses of the five samples for the 88u entity (pentadecanoic acid, ethyl ester), showing its presence only in samples B and D.

### Conclusions

This approach successfully generated chemometric profiles that resulted in the classification of the five whiskey samples into four groups, and could additionally be used to analyze for unintended contamination [3], optimization of product storage conditions, and determination of sample deterioration over time.

### References

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- L. Vaclavik, , O. Lacina, J. Hajslova, J. Zweigenbaum, "The use of high performance liquid chromatographyquadrupole time-of-flight mass spectrometry coupled to advanced data mining and chemometric tools for discrimination and classification of red wines according to their variety.", Anal Chim Acta. 685, 45-51 (2011).
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