Metropolitan Museum of Art Gas Chromatography- Mass Spectrometry (GC-MS) Results from Material Analysis

This document includes (1) a mass spectrum and (2) the volatile organic compounds (VOCs) emitted from samples using GC-MS analysis. The data is not interpreted; however, several classes of chemicals are highlighted because they are potential risks for artwork in an enclosed environment. A basic key, provided below, indicates those classes. The amount of each chemical identified has not been determined; similarly, it is not known how much of each chemical is necessary to do damage to art. Finally, peaks may be present that are the result of the sample adsorbing chemicals from the air and reemitting them during testing rather than being inherent to the sample. Research is ongoing to determine specifically which chemicals and amounts are required to negatively affect artifacts.

Highlighted data:

Pink – chemicals currently known to be hazardous to art

Green – amines; can raise the pH, are suspected to react with acids and may form crystals in an enclosed environment

Yellow – chemicals of the following type, which may be hazardous to art:

Acids – lower the pH, corrosive to metals, degrade organic materials

Aldehydes – can convert to acids with heat or exposure to UV light

Esters – can hydrolyze into acids with heat and humidity

Sulfur-containing compounds – known to tarnish and corrode some metals

Halogenated compounds – can become reactive with exposure to heat and UV light

Nitrogen-containing, not amine – can react with other off-gassed chemicals

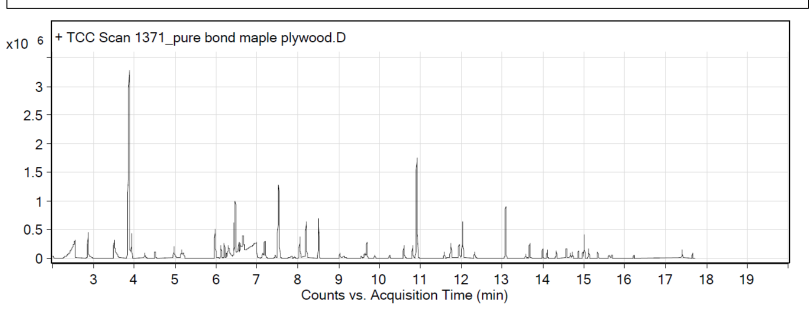
Alkynes – can become reactive when exposed to heat or UV light

Sample: Purebond maple plywood; Manufactured by: Columbia Forest Products

Oddy test score: Unsuitable Date collected: 08/15/2016

Technique used: SPME with a PDMS/DVB fiber; Agilent 7890B GC and 5977B MS fitted with a GL Sciences OPTIC-4 multimode inlet and LEAP PAL RTC autosampler; Pre-heated at 60°C for 20 minutes; fiber exposure at 60°C for 20 minutes; sample injected into 220°C inlet and crotrapped for 2 min at -15°C; GC ramped from 40°C to 225 °C at 10°C/min. Data analyzed in masshunter Qualitative. Samples > 90% match with a NIST 17.0 library are reported.

VOCs not highlighted are because they were also observed in blanks: (1) 12.0 min: 2-methyl-, 3-hydroxyl-2,4,4-trimethylpentyl ester propanoic acid



RT	Mass	Name	DB Formula
2.55		Acetic acid	C2H4O2
2.87		Pentanal	C5H10O
3.51		1-Pentanol	C5H12O
3.87		Hexanal	C6H12O
3.94		Cyclotrisiloxane, hexamethyl-	C6H18O3Si3
4.25		2-Furancarboxaldehyde	C5H4O2
4.97		2-Heptanone	C7H14O
5.98		2-Heptenal, (E)-	C7H12O
6.12		Benzaldehyde	C7H6O
6.2		Cyclotetrasiloxane, octamethyl-	C8H24O4Si4
6.47		Furan, 2-pentyl-	C9H14O
7.19		3-Octen-2-one	C8H14O
7.53		2 OCTENAL	C8H14O
7.55		(S)-2-Methyl-5-oxotetrahydrofuran-2-carboxylic acid	C6H8O4
8.2		Nonanal	C9H18O
9.03		2-Nonenal, (E)-	C9H16O
9.69		Decanal	C10H20O
10.59		6-Undecanone	C11H22O
10.81		Hexanoic acid, pentyl ester	C11H22O2
11.59		5-pentyl-2(5H)-furanone	C10H18O
11.95		6-Dodecanone	C12H24O
12.04		Propanoic acid, 2-methyl-, 3-hydroxy-2,4,4- trimethylpentyl ester	C12H24O3
14.1		Benzene, (1-propylheptyl)-	C16H26
14.32		Benzene, (1-ethyloctyl)-	C16H26
14.66		Hexadecane	C16H34
17.4		Hexadecanoic acid, methyl ester	C17H34O2
17.65		1,2-Benzenedicarboxylic acid, dibutyl ester	C16H22O4