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Testing For Free (Unbound) Sulfur Dioxide In Wine Using The Aeration Oxidation Method

This instruction sheet also includes very good information on how to care for and prepare laboratory equipment to get the most accurate results

Categories of Sulfur Dioxide (hereafter called SO₂): SO₂ is added to wine as a preservative and protector.

- Free (**unbound**) SO₂ – It is this free SO₂ that is the ‘workhorse’ at protecting wine.
- Bound SO₂ – This can be measured but it is not as critical that it be done. Some of the bound SO₂ is loosely bound and can release into free SO₂ when running some tests. Care must be taken to prevent this in order to avoid a false reading.
- Total SO₂ – This is merely the sum of the bound SO₂ and the free SO₂.

For many Aeration Oxidation is the most reasonably accurate method to test SO₂ levels in wine. There is a slight overestimation of the SO₂ level in red wine because some of the SO₂ that is bound to color pigments will release and be erroneously read as being free SO₂. The older the red wine is the less this occurs. At any rate the overestimation is slight and not of significant consequence.

Labware Needed:

I. Aeration Oxidation apparatus for measuring free SO₂

A. The parts include: *(if more than one is needed that number is shown in ()):*

- Metal stand (**rod & base**)(*there may be some value in having two stands*)
- 100 ml round bottom double neck flask
- Clamp (**to attach the 100 ml flask to the metal rod**)
- Bored rubber stopper # 5 with a ¼ inch hole
- Bored rubber stopper # 5 with a ⅜ inch hole
- Pasteur pipet (**2**) (*Only one is needed but an extra is included because of fragility*)
- Glass tip
- Tubing to run from the top of the 100 ml flask to the upright on the impinger top
- Complete impinger unit (*this means it has a top & a bottom*).
- Clamp (**to attach the impinger bottom to the metal rod**)
- Tubing to run from the side attachment of the impinger top to the inlet (**bottom**) of the flowmeter
- Flowmeter
- Tubing to run from the outlet (top) of the flowmeter to the water aspirator (*or a catch basin if an aqua pump is used*)
- A plastic quick disconnect to insert somewhere inline on the tubing that runs from the aspirator to the impinger (*this allows for disconnecting the A/O apparatus from the water aspirator*).
- Plastic tie clamps (**2**) (*to secure the flowmeter to the metal rod on the stand*)
- Water aspirator (*an aqua pump can be substituted with 1200 being a good size*)
- Sink adaptors (**2**) (*to accommodate attaching the aspirator to the sink*)
- **Note:** To test for bound SO₂, the addition of a condenser, a heating element and a longer metal rod would be required. These may be special ordered. An aqua pump cannot be used when testing for bound SO₂. The water aspirator must be used because water must be drawn through the condenser to cool the sample during testing.

Note: The recommended number of the labware listed below can possibly be reduced if there is a willingness to clean a single piece before using it to handle a different substance.

- II. 10 ml volumetric pipet (**2**)
- III. 20 ml volumetric pipet
- IV. Pipet bulb (used to draw fluid into a pipet) (adding a glass tip to the bulb increases ease of use)
- V. Small beakers (**3**)
- VI. 10 ml or 25 ml buret (**2**) with stopcock (*a smaller volume buret simply may require refilling*)
- VII. Buret stand with buret clamp

- VIII. Some type of ice bath container in which to suspend the 100 ml double neck flask (*Tupperware works well*). During the test, keep the wine sample temperature below 20° C (68° F)
- IX. 100 or 200 ml volumetric flask (3) These are used to prepare and/or hold reagents
- X. 250 or 500 ml plastic rinse bottle with rinse spout (for distilled water)
- XI. Safety glasses (*recommended when using chemical reagents*)
- XII. A timing device (*having a second hand is desirable & a stopwatch would be ideal*)
- XIII. Grease pencil with which to label beakers, pipets and etc.

Reagents Needed: These can be purchased or made from stronger solutions by proper dilution using distilled water. Instructions for some dilution procedures are included with this document.

- XIV. 0.3% hydrogen peroxide (*hydrogen peroxide should be stored in a refrigerator as it is unstable at room temperature*). (*We have come across some controversy concerning the strength of hydrogen peroxide needed. Virtually all the literature calls for using 0.3% but a few winemakers have said they can only get accurate results by using 3.0% (the drugstore kind) hydrogen peroxide. Numerous attempts on our part to clarify this have thus far proven futile.*)
- XV. 25% (3 + 1) phosphoric acid
- XVI. SO₂ indicator solution (*made from methyl red and methylene blue*)
- XVII. 0.01 normal (0.01N) sodium hydroxide (NaOH)
- XVIII. Distilled water

Procedures: There are many steps but don't panic, they are quite easy.

Step One: Consider labeling the beakers, pipets, flasks and etc. with a grease pencil as to what reagents they will contain. Label a large beaker as a waste beaker in which to dump waste reagents. **It is recommended that safety glasses be worn when working with chemicals. Make sure all labware is clean.**

Step Two: Set up at least one buret on its stand. Pour a small amount of 0.01 (0.01N) normal sodium hydroxide (NaOH) into a small beaker, swirl to give the beaker a final cleansing and discard into the waste beaker. Then pour an adequate amount of the 0.01N sodium hydroxide into the beaker, which then can be used to pour the reagent into the buret. Pour a small amount of the sodium hydroxide into the buret, tilt the buret on its side with the open end held over the waste beaker. Roll the rinse solution around in the buret and then discard the rinse into the waste beaker. Pour, from the beaker, an adequate amount of fill into the buret. A small buret funnel can be very helpful in pouring liquid into the buret. **It is not absolutely necessary to fill the buret completely to the zero level at the top but be sure to have enough for the needed task.** You should dispense liquid out of the buret into the waste beaker until all air bubbles are eliminated. You can possibly hold off on setting up a 2nd buret and certainly hold off on pouring in hydrochloric (*or other acceptable*) acid until such time as it really is needed. Use the same procedure for adding hydrochloric acid to the 2nd buret as was used for the sodium hydroxide. These burets can now be set aside for use later. Using a buret to dispense a liquid is known as titration.

Step Three: If not already done, assemble the aeration oxidation apparatus (*see attached schematic*).

Step Four: Use a plastic rinse bottle of distilled water to rinse out the 100 ml round bottom double neck flask. Discard the water into the waste beaker or sink. Clamp the top neck of the flask to the metal rod and suspend it in the ice bath to begin cooling. Nothing will be added to the flask just yet. Insert the heavier but shorter glass tip into the # 5 rubber bung with the ³/₈ hole and insert into the top neck of the flask. Attach the proper tubing to the protruding tip and attach the other end of the tubing to the impinger top. **The ice bath will be most effective and easiest to use if it contains mostly cold water with some ice.** It is important that the flask be cold when the wine sample is added to reduce the chance of some of the loosely bound SO₂ becoming free SO₂.

Step Five: Make sure the collecting impinger is clean & dry. Prepare a small beaker and a 10 ml volumetric pipet to receive an adequate amount of 0.3% hydrogen peroxide (*as mentioned above, a few use 3.0%*). In a similar fashion as was done for the sodium hydroxide above, give a final cleansing rinse to the beaker and the pipet. The pipet should be tilted on its side and the rinse rolled around as was done in the preparation of the buret. Discard this rinse into the waste beaker. Using a pipet bulb, draw the hydrogen peroxide solution to fill a 10 ml volumetric pipet. Draw the liquid up past the fill line, pull off the bulb and quickly put your finger on the top of the pipet to prevent premature drainage of the solution. **It is really helpful if your finger is very dry when holding and later releasing the liquid from the pipet.** Release the excess peroxide into the waste beaker being very careful not to allow the liquid to fall below the etched gradation line. Read the amount at the bottom of the meniscus. Also it is much better to raise the pipet up to eye level to get a good reading rather than stooping down to get the reading. Release the 10 ml of hydrogen peroxide into the bottom part of the impinger unit.

Step Six: Add three drops of the SO₂ indicator solution into the impinger bottom. Swirl to mix it into the already present hydrogen peroxide. Be careful to avoid getting the indicator anyplace other than into the impinger as its stain is dark and will linger long. You should get an olive green or turquoise green color in this mixture that is in the impinger. If you have that color at the start, there will be no need to adjust it. However, if you don't have that color, then you can adjust it by first adding a drop of the 0.01N sodium hydroxide (*some even use 0.1N for this procedure*) into the buret you may have prepared earlier followed, if needed, by a drop of mild hydrochloric acid (*or suitable*)

substitute) from another buret. Toggle back and forth between the two until you get the required olive or turquoise green color. The sodium hydroxide will have a greening effect and the hydrochloric (*or other*) acid will have a purpling effect. **Remember, if you had that required olive or turquoise green color from the start you would not need to titrate with the sodium hydroxide or the acid.** (*And to repeat again, it may be possible to substitute a different kind of mild acid such as mild phosphoric or mild sulfuric acid as well as a different normality of NaOH as the purpose is merely to adjust the color*).

Step Seven: Clamp the impinger bottom to the metal rod. Then the impinger top can be inserted into the impinger bottom. Setting up the impinger with its solution should be done first before the double neck flask is prepared. This will increase the chance of still capturing any SO₂ that may blow off prematurely when the wine sample is added to the 100 ml flask.

Step Eight: Prepare, by rinsing with 25% phosphoric acid, a beaker and 10 ml volumetric pipet using the same techniques as described previously. Then draw a 10 ml amount of 25% phosphoric to then be released into the double neck flask via the side neck. It is important to add the phosphoric acid to the flask before the wine sample is added to reduce the chance of any premature loss of any of the free SO₂ for which you are testing. The phosphoric acid will make the wine more acidic (lowering the pH) so it more easily will release the free sulfur dioxide (SO₂).

Step Nine: From the wine sample you hopefully collected in advance, you will now measure out the correct amount (**20 ml**) to be added to the double neck flask. The wine to be tested should have been stored in an inert (**glass**) container that is full to the brim. This will reduce the risk of prematurely losing any free SO₂. Again, as done numerous times above, rinse a beaker and 20 ml volumetric pipet with a small amount of the wine and discard into the waste beaker. Pour an adequate amount of wine **down the side** of the beaker. Then draw the 20 ml sample of wine and release it into the double neck flask via the side neck. Use the ice bath to keep the sample below 20^o C (**68^o F**). Stopper the side neck with the # 5 rubber stopper with the ¼ inch hole. Prior to this slip the long, thin and very delicate Pasteur pipet into the stopper hole (*lubricate the pipet with Antifoam or glycerine to ease insertion of the glass tube*). Position the bottom tip of this pipet so it rests into the wine/phosphoric acid mixture inside the flask. This will allow the air to bubble through the sample.

Step Ten: It is now time to aerate the wine sample. This can be done one of two ways. Either hook up a vacuum pump to the Pasteur pipet protruding from the side neck of the double neck flask to push air through the system or attach the aspirator to a sink faucet and then connect the other end of the tubing to the flowmeter outlet (**top**) in order to pull air through the system. The aeration should be at a rate of 1000 to 1500 ml/minute (**1.0 to 1.5 liters per minute**) as measured by the flowmeter. Run the aeration for ten (**10**) minutes at which time the airflow is shut off. The collecting of the free SO₂ in the impinger causes a reaction with the hydrogen peroxide resulting in a conversion to sulfuric acid which can then be titrated as discussed in steps 11 & 12 below.

Step Eleven: Now it is time to analyze the mixture collected in the impinger. Pull the tubing off the top of the impinger and lift the top partially out of the bottom container. While keeping the tip of the top's glass tube inside the bottom, use the rinse bottle of distilled water to rinse off any of the purple collection still clinging to the outside circumference of the tube. Also squirt some water down inside the impinger top to push any collection back into the liquid in the bottom. **Distilled water will not have any affect on the collection material.** Now the impinger bottom can be dismantled from the apparatus. The contents will be a purple color. Using the buret containing 0.01N sodium hydroxide (**NaOH**), titrate carefully (**drop by drop until you get more adept**) the contents of the impinger. It is vital that you first record (**yes write it down**) an initial reading of the level of sodium hydroxide contained in the buret. **Swirl the impinger contents at each addition of NaOH.** Titrate until the purple color changes back to its original olive or turquoise green color. It is imperative that the exact last drop that permanently changes the color from purple to the olive or turquoise green is observed and the titration stopped immediately. If titration takes you past the olive or turquoise green color, a false reading will occur and the whole test will require repeating. You know you went past the end point if the color changes to a lighter & brighter green. The further past you go the lighter will be the color. **A sign that the endpoint is very close is that the purple will start to turn dingy with a gray hue followed by a gray color. The next drop of sodium hydroxide may indeed cause the transition back to the original olive or turquoise green color so definitely go drop by drop.** After reaching the proper end point, read the level of sodium hydroxide left in the buret. Subtract the initial reading from the final reading to know how much sodium hydroxide it took to get back to the original color. **Record these numbers to avoid error.**

Step Twelve: Calculate the free SO₂ level using the following formula:

$$\text{Free SO}_2 = \frac{(V) * (N) * (32) * (1000)}{(v^1)}$$

Where: (**V**) =The amount of sodium hydroxide used to titrate back to the original color (*this variable is the only truly unknown one and won't be known until you actually do the titration*).

(**N**) = The normality level of the sodium hydroxide used (*you know this really before you start and unless you change to a different normality it will cease to be an unknown variable*).

(**v¹**) = The size of the wine sample (*if you followed these directions the wine sample was 20 ml so that also ceases to be an unknown variable*).

Therefore: You can work out most of the formula ahead of time to be: **Free SO₂ = (V) * 16**

For Example: If it took 2.85 ml of the 0.01N sodium hydroxide to turn the purple color back to the proper green color, the free sulfur dioxide level would be 2.85 * 16 = 45.6 parts per million (ppm) sulfur dioxide.

Procedures For Making Various Reagents

Hydrogen Peroxide 0.3 %:

- Use one milliliter (*ml*) of 30% hydrogen peroxide and then add distilled water to make a solution that totals 100 ml. Use a 100 ml volumetric flask to make one batch or a 200 ml volumetric flask to make a double batch. 30% hydrogen peroxide should be readily available from winemaking supply stores.
- An alternative is to use 10 ml of 3% hydrogen peroxide and bring to a 100 ml volume by adding distilled water. Again, use either a 100 ml or 200 ml volumetric flask. 3% hydrogen peroxide is readily available from your local drug store.
- Hydrogen peroxide decomposes fairly quickly at room temperature so store the 30% or 3% solutions in the refrigerator until needed. It is best to make up the 0.3% solution as fresh as possible before use.

Phosphoric Acid 25% (3 + 1):

- Add 294 ml of 85% phosphoric acid to a one-liter flask. Add distilled water to bring to the one-liter volume. 25% solution already made up is available from winemaking supply stores.

Sulfur Dioxide Indicator:

- Into 100 ml of 50% ethyl alcohol, dissolve 0.1 gram of methyl red and 0.05 gram of methylene blue.
- Ready made solution is available in small quantities from supply stores.

Sodium Hydroxide (NaOH) 0.01 Normal: Also written as N/100 or 0.01 N or 1/100N.

- Put 10 ml of 1/10 Normal (**0.1N**) NaOH into a 100 ml volumetric flask and add distilled water to bring the flask to volume. If the flask is a 200 ml flask use 20 ml of the NaOH and distilled water to make volume. One-tenth normal NaOH is very readily available from winemaking supply stores. It is the most commonly used strength for doing the common T/A acid test in wine.
- An alternative is to make the solution from one normal (**1.0N**) NaOH. Add 10 ml of the 1.0N NaOH to a 1-liter (**1000 ml**) volumetric flask. Add distilled water to make volume in the flask.
- NaOH can stray from its intended normality level especially with higher exposure to oxygen in the air. There are chemicals (**potassium acid phthalate or hydrochloric acid & methyl red**) available from winemaking supply stores that can be used to recalibrate the normality of NaOH. Or, fairly regularly repurchase a new batch of NaOH.

Following Are Two Formulas That May Be Of Value:

For Normality - Amount To Add = $\frac{\text{Normality Desired} * \text{Volume Desired}}{\text{Normality of the Concentrated Solution}}$

For Example – $\frac{0.01N * 100 \text{ ml}}{0.1N}$ = 10 ml of 0.1N NaOH added to a 100 ml flask, brought to volume with distilled water will make 0.01N NaOH.

For Percentage – Volume To Add = $\frac{\% \text{ Desired} * \text{Total Volume Desired}}{\% \text{ Of The Concentrated solution}}$

For Example – $\frac{0.3\% * 100 \text{ ml}}{30\%}$ = 1 ml 30% hydrogen peroxide added to a 100 ml flask, brought to volume with distilled water will make 0.3% hydrogen peroxide.