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# SURFACE COLOUR CHANGE IN WOOD DURING DRYING ABOVE AN BELOW FIBRE SATURATION POINT

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#### **ABSTRACT**

A technique, useful for studying the formation of kiln brown stain in wood drying, has been develor to measure the surface colour change in a single board wood sample during drying. The wood sam is planed carefully in the green state to remove any surface wood that was damaged during cutting. intact tracheids at the surface cause the evaporative front to remain at the surface during drying therefore colour formation also occurs right at the surface. In this way, the colour can be measured us a spectrophotometer at various stages during drying without having to slice the sample.

Experiments were carried out to measure the change in colour of wood from green to E corresponding to the drying schedule used. At the end of each schedule the boards were held at EMC to determine how the colour changed below fibre saturation point. The results show that colour of the wood continues to change below the fibre saturation point and the nature of the co change indicates an increase in the complexity of the coloured compounds present.

Further experiments were done to measure the rate of colour development at different temperatusing the technique developed. The results have shown a correlation between temperature and co development over the range 50°C to 70°C with the rate increasing significantly above 60°C.

**Keywords:** kiln brown stain, wood colour, colour measurement, wood drying.

### INTRODUCTION

*Pinus radiata* is known for its bright colour, clear ring pattern and medium strength which mal suitable to a wide variety of end uses such as panelling, mouldings, joinery and furniture. Howe wood colour changes during drying when high temperature schedules are employed which adver affects its reputation. This colour change can be in the form of an overall change in colour or formation of kiln brown stain (McCurdy *et. al.*, 2003). Kiln brown stain has the greater effect on qua for high value appearance products. This is an irregular brown colouration that occurs 1 to 2 mm the surface of drying boards that is considerably darker than the surrounding wood. This layer is consist with the thin dry layer formed in the kiln drying of softwood lumber (Pang *et al.*, 1994).

Studies by McDonald *et al* (2000) have shown that kiln brown stain in *Pinus radiata* is most like caused by a Maillard reaction between sugars and amino acids in the wood sap as it is concentrate the wood surface during drying.

Various methods have been developed to measure kiln brown stain formation in samples of w during drying. Studies by Kreber and Haslett (1997) used visual inspection to determine the leve stain whereas Dawson *et. al.* (2003) used a colorimeter for the same purpose. More recently, Ledig Seyfarth (2001) have used a spectrophotometer to measure surface colour in European beech and h successfully characterised the wood colour using the CIELab system. Photographic techniques (McCu et.al., 2001; Dieste, 2002) and microscopic techniques (McCurdy et.al., 2005) have also been develothat combine digital photography with computer image analysis.

The major difficulty encountered when measuring kiln brown stain is that the most significant is formation occurs about 1mm below the surface of the timber. This means that any measurement of surface may not measure the full extent of the stain formed. Also if the surface is removed to reveal stain it is a destructive test so cannot be performed while the sample is being dried in the kiln. The paper describes a technique developed to overcome this problem and results from initial experiment determine colour change rates will be discussed.

### **MATERIALS AND METHODS**

The samples used were cut from green *Pinus radiata* boards to a size of 25x25x100mm. The ends two of the sides of the samples were sealed with paint so that only the top and bottom flat faces (tangen were available for evaporation of moisture. Then, the samples were stored in a freezer until further proces for the drying tests.

Immediately before drying, the surfaces of the samples were prepared by carefully smoothing them a hand plane. The objective of this step was to remove the damaged layer from the surface of the wood ge so that there are intact tracheids near the surface. The advantage of planing the surface is that the blade sl through the wood which gives a smoother finish and causes less damage to the cells on the surface.

The surface to be measured was marked at sampling areas before drying commenced so that the part the surface measured were consistent the same throughout the drying process. This was done by mark each point with a circle slightly larger than the aperture of the spectrophotometer. In the experiments conduct four measurement points were used for each sample.

The measurement of the surface colour during drying was performed by removing the sample from dryer, recording the time and weight. The colour of each of the measurement points was then determ using a surface reflectance spectrophotometer (Minolta CM-2500d portable integrating spl spectrophotometer). This instrument uses two pulsed xenon lamps as a light source, a diffraction grating spectral separation and a silicon photodiode array as the light-receiving element. This array is a dual element device that can measure at 10nm intervals over the range from 400nm to 700nm.

In addition a photograph was also taken of the surface each time when the colour was measured qualitative analysis of the colour development. The entire measurement process only took a few minuthus keeping the cooling and drying of the sample outside the dryer to a minimum.

The results presented in this paper are from seven drying runs, the first three of which were performe test the technique and determine if colour development changes below fibre saturation point. The remain four drying runs, and one of the previous runs (initial run 3 is the same as rate run 5) were used to determ the rate of colour development during drying at different drying temperatures. In each of the drying runs

**Table 1.** Drying schedules and holding conditions for the eight drying runs.

Run No.	Drying conditions			Holding conditions		
	Dry bulb	Wet bulb	Elapsed time (h)	Dry bulb	Wet bulb	Elapsed time (h)
Initial Run 1	50°C	40°C	48	50°C	48°C	50
Initial Run 2	70°C	60°C	48	70°C	69°C	26
Initial Run 3*	50°C	40°C	75	70°C	69°C	48
Rate Run 1	50°C	30°C	48	50°C	49°C	36
Rate Run 2	50°C	30°C	48	55°C	54°C	36
Rate Run 3	50°C	30°C	48	60°C	59°C	36
Rate Run 4	50°C	30°C	48	65°C	64°C	36
Rate Run 5*	50°C	40°C	75	70°C	69°C	48

<sup>\*</sup> These are the same run.

## RESULTS AND DISCUSSION

The spectrophotometer measurements produced a spectrum for each area measured at different dry times as well as the colour expressed using the CIELab colour coordinates  $L^*$ ,  $a^*$  and  $b^*$ . From colour coordinates, the colour change was then calculated ( $\Delta E$ ) using the following equation, where the overall change in colour:

$$\Delta E = \sqrt{\Delta L^2 + \Delta a^2 + \Delta b^2}$$

The time point with the greatest value for lightness was chosen as the datum with which the chain wood colour was compared. This point occurred within a few hours of drying when the surface the samples had dried out but discolouration had not begun. The  $\Delta E$  values calculated for each run been plotted in Figure 1 to Figure 3, with the moisture content also plotted on these graphs.

The results for the Initial Run 1 (Figure 1) show that the colour change drops sharply at the beginn of drying, as the wood colour gets lighter due to the surface losing moisture. After reaching plightness at about 5h into the schedule the surface colour of the sample darkens in an approximal linear relationship with elapsed time. There appears to be a slight change in the slope at 72h when schedule was changed to holding conditions to maintain the moisture content.

The results for the Initial Run 2 (Figure 2) show a similar increase in lightness at the beginning

The results for the Initial Run 3 (Figure 3) also show a sharp increase in lightness when the surbegins drying. This is followed by a slow increase in colour, up to the point where the schedul changed to the holding conditions at 78 hours. At this point the colour change increases sharply then gradually decreases when the colour change appears to tend toward an asymptotic value in a sim way to the Initial Run 2.

The subtraction spectra for the Initial Run 3 are plotted in Figure 4. These spectra were obtained subtracting the 4h spectra from the spectra of each measurement after 4h. This means that the higher absorbance value of light at a particular wavelength the darker the colour. The spectra show that drying progresses there is an overall increase in light absorption across all wavelengths. As dryprogresses there was also a shift to a higher wavelength of peak (red shift).

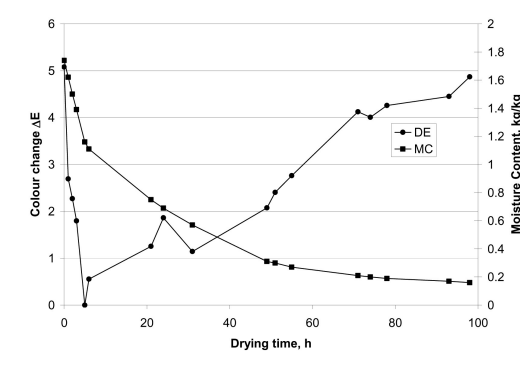
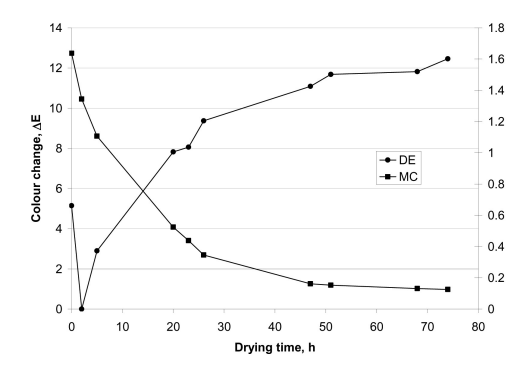
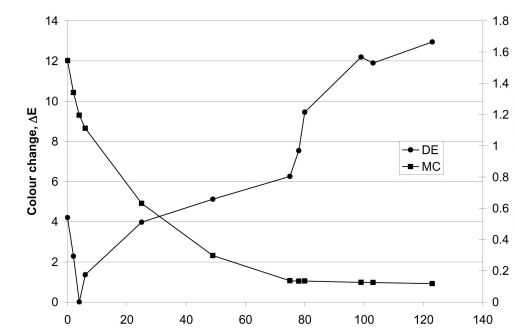


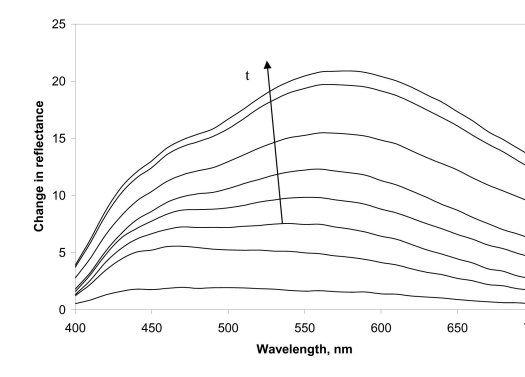
Figure 1. Results for Initial Run 1 showing the change in colour ( $\Delta E$ ) and moisture content (MC a function of elapsed time.





**Figure 2.** Results for Initial Run 2 showing the change in colour ( $\Delta E$ ) and moisture content (MC) a function of elapsed time.

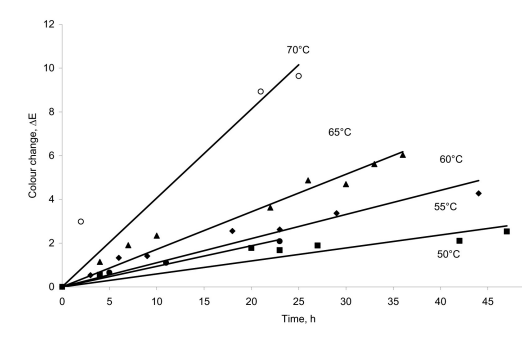




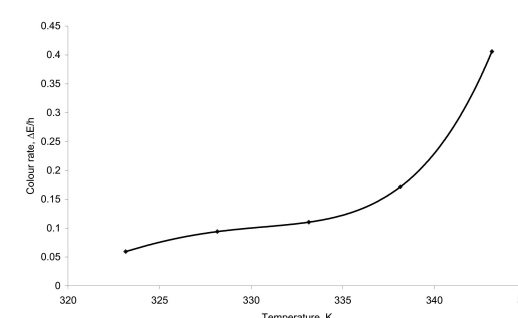
**Figure4.** Subtraction spectra for Initial Run 3 with the values showing spectra of each measurement after 4 hours being subtracted by the spectrum at 4 hours.

The results for the rate determination experiments (Rate Runs 1 -5) are arranged to the colour change in the approximately linear region as a function of elapsed time as sho in Figure 5. These data are generally a good linear fit except for the 70°C data (too few opoints).

Most importantly, the rate of colour change clearly shows an increase with dry temperature. In order to quantify the relationship between the temperature of the dry schedule and the rate of colour changes, the colour change rates for all of the runs are plo in Figure 6. It has been found that such relationship was non-linear for the data produced this experiment and could only be fitted by a 4<sup>th</sup> order polynomial. This may be an error the experiment or may be an accurate representation of colour change in wood. The shows that the colour change rate increases remarkably between 65°C (338K) and 7 (343K). The data could not be fitted using the Arrhenius equation.



**Figure 5.** Colour change with time for 5 different drying temperatures.



From the results described above, it was observed that in the (Rate Run 1 and 2) Initial Rudrying run at first seem to suggest that there was a reduction in the rate of colour development moisture content decreases. The results from the Initial 3 run however show that the higher of colour development caused by the increase in temperature from the drying schedule to holding schedule was not affected by the moisture content. This means that there must be ano explanation for the change in rate of colour development.

The alternative explanation is that there is a maximum possible colour development caused initial reaction of sap compounds based on the concentration of these compounds in the before drying. Once these compounds have reacted, further reactions become more complex proceed at a slower rate. The early stages of the Maillard reaction are reasonably well understout beyond the first (Amadori) products there are a large number of possible reactions (Fayle Gerrard, 2002a), so it is difficult to imagine the actual structure of the coloured compound produced.

The shift towards more complex reactions is supported by the shift to higher wavelength the subtraction spectra. The colour exhibited by the Maillard reaction products is most like caused by a combination of extended polyene structures attached to electron-donor and –accel groups as is the case with most organic colorants (Nassau, 2001). The addition of more conjugated double bonds or donor-acceptor groups will cause the reaction products to absorb lon wavelengths. Studies of the Maillard reaction have shown that complex products containing cypolyenes and donor-acceptor groups are produced (Fayle and Gerrard, 2002b).

The development of colour below fibre saturation point at a similar rate to colour developm above fibre saturation point suggests that the majority of colour development is occurring in cell walls. This is because the only water in the wood below fibre saturation point is the bo water in the cell walls. Digital micrographs of stain layers do show that the majority of brown stain is due to darkened cell walls (McCurdy et. al., 2005).

The rate measurement experiments show clearly that there is a non-linear relationship betw temperature and the rate of colour development. This is to be expected for any chemical react though in this case it was not possible to fit the data to the Arrhenius equation. This is likely to due to there being more than one reaction involved in the formation of colour. Regardless of reaction mechanism the rate of colour development increases greatly above 60°C.

The experiments have shown that the smoothing of the surfaces of green wood samples wi plane has caused the kiln brown stain to form closer to the surface where it has been successf measured using a reflectance spectrophotometer.

In normal rough-sawn timber the evaporative front recedes through the layer of dama tracheids at the surface early in drying forming what is known as the thin dry layer (McCurdy Keey, 2002). This means that most colour development occurs at 1 to 2mm below the surface is not always apparent on the surface. The process of removing the damaged layer means that evaporative front stays at the surface and the formation of colour is more obvious and car measured non-destructively during the drying process.

This technique for measuring the development of colour in wood during drying shows potent to be a useful research tool and results from initial experiments show that further research

### **CONCLUSIONS**

The results presented here have shown that surface colour formation can be measured during if the surface is planed in the green state. Using this method the experiments perform have measured the rate of colour formation between 50°C and 70°C and shown that the rate greatly increased above 60°C. The experiments have also shown that colour formation occubed above and below the fibre saturation and the moisture content has little effect on the rate colour formation.

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