Comparative studies on the mechanical properties and microstructures of outerwood and corewood in *Pinus radiata* D. Don

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Abstract

Twenty-year-old *Pinus radiata* trees imported from New Zealand were investigated, and a comparison was made between the outerwood (rings 16–20) and corewood (rings 4–6) in terms of mechanical properties, anatomical characteristics, microfibril angle (MFA), relative crystallinity, crystallite size and lignin content to determine the relationship between their mechanical properties and microstructures. The results demonstrated that the mechanical properties of the *Pinus radiata* outerwood were significantly better than those of the corewood. The outerwood had a modulus of rupture (MOR) of 106 MPa, a modulus of elasticity (MOE) of 11.4 GPa, and compressive strength parallel to the grain of 38.7 MPa, and the MOR, MOE and compressive strength parallel to the grain of the corewood were 78.9 MPa, 7.12 GPa and 29.3 MPa, respectively. The observed microstructures of the *Pinus radiata* outerwood and corewood were different, mainly due to differences in cell wall thickness, MFA, and relative crystallinity. The double wall thickness of the tracheid cells of the corewood and outerwood were 3.65 and 5.02 µm, respectively. The MFA data indicated that the average MFA of the outerwood was 14.0°, which was smaller than that of the corewood (22.3°). With X-ray diffraction, the relative crystallinity of the corewood was determined to be 35.7%, while that of the outerwood was 40.2%. However, the crystallite size of the outerwood cell wall shows no obvious difference from that of the corewood. Imaging FTIR spectroscopy showed a slightly higher relative content of lignin in the cell wall of the outerwood. The correlation between the microstructures and mechanical properties showed that the corewood with a thin cell wall, large MFA and low relative crystallinity had poor mechanical properties, while the outerwood with a thicker tracheid, smaller MFA and higher relative crystallinity had better mechanical properties. This means that the MFA, relative crystallinity and cell wall thickness synergically affect the mechanical properties of *Pinus radiata* in different radial locations.

Keywords: *Pinus radiata*, Outerwood, Corewood, Mechanical properties, Microstructures

Introduction

China is a country that engages in the export and consumption of wood products on a large scale, but it is also a country with relatively scarce forest resources. In recent years, with the increase in domestic timber cutting restrictions, China’s timber dependence on foreign countries has increased significantly [1]. *Pinus radiata* is the main afforestation tree species in New Zealand, which has the characteristics of rapid growth, straight wood grain, uniform structure, and strong stability, and is one of the most imported wood species in China in recent years [2]. Thus, research on the properties and processing technology of *Pinus radiata* has become one of the hotspots in the wood academic field.

There have been some studies conducted on imported *Pinus radiata*, which have mainly focused on processing and utilization [2], heat treatment modification [3] and the physical and mechanical properties [4]. Imported *Pinus radiata* usually has a large diameter, with an average diameter of 46 cm for trees 20 years old.
in previous research, and radial variations in the physical and mechanical properties of *Pinus radiata* have been preliminarily investigated [4, 5]. For softwood, the concepts of juvenile versus mature wood, or corewood versus outerwood are usually used to discuss the radial variation in such properties [6–8]. Research on 17-year-old *Pinus radiata* from Chile and 11-year-old *Pinus radiata* from New Zealand showed radial variations in the modulus of elasticity (MOE), microfibril angle (MFA) and cell wall thickness [9, 10]. Research on the properties of 27-year-old *Pinus radiata* from New Zealand showed that the density and MFA are significantly different between juvenile wood and all-wood composite [11]. For 13-year-old *Pinus radiata* from Australia there is a significant relation between MOE and age [12]. However, the mechanical properties and microstructures of different radial locations and their correlation in *Pinus radiata* have not been systematically reported.

The cell wall is the solid material of wood and the bearing structure of wood under stress. The anatomical structure, ultrastructure and the chemical composition of the cell wall affect the wood properties [6, 13, 14]. Studies have shown that the variation in the cell wall thickness may become an important factor when analyzing the mechanical behavior of a wood cell [7]. As one of the important ultrastructural factors of the cell wall, the effect of MFA on the mechanical properties of wood is an important research topic. The change in MFA was found to be negatively correlated with the change in MOE, modulus of rupture (MOR) and hardness [9–11].

Different from the MFA, the crystallinity of the wood cell wall is positively correlated with the elastic modulus of wood. It was found that the decrease in the MOE of Chinese fir is related to the decrease in the crystallinity of the wood cellulose [15]. The correlation coefficient between the *Populus tomentosa* cell wall crystallinity and MOE was 0.75 [16]. In the model for predicting wood hardness, when the latter quantifier of crystallinity was not involved, the width and length of crystallites were consistently positively related to tangential and radial hardness but negatively related to transverse hardness [17], while the effect of crystallite size on other mechanical properties has not been reported.

As the substrate material of the cell wall, lignin is one of the factors that affects the elastic modulus of the cell wall, and lignin content is positively correlated with the elastic modulus of the cell wall [18, 19]. In Gindl’s research, the longitudinal modulus of elasticity was higher after the completion of lignification in Norway spruce [20], which explained why the MOE of the secondary wall of mature tracheids with high lignin content in spruce was 22% higher than that of immature tracheids [21]. In summary, the microstructures of wood have a significant impact on its mechanical properties, but the microstructural characteristics of 20-year-old *Pinus radiata* at different radial positions and how these structural characteristics affect the mechanical properties have not been reported.

Research on the differences between corewood and outerwood at the cell wall level is becoming important because it can clarify the macroscopic differences in the wood mechanics. The purpose of this study was: (1) to provide mechanical data, and microstructure characteristics and compare the differences between corewood and outerwood in 20-year-old *Pinus radiata* imported from New Zealand, and (2) to analyze the effect of microstructures on their mechanical properties through the revealed results from (1).

### Materials and methods

#### Materials

Logs (5.9 m) of *Pinus radiata* imported from New Zealand at the age of 20 were selected as experimental materials. Bolts 1.5 m in length were taken from six different logs for the measurements. A total of 6 bolts were measured. The sampling method is shown in Fig. 1. Two corewood (rings 4–6) and two outerwood (rings 16–20) strips were taken from each bolt. A total of 24 samples from the corewood and outerwood were used to perform the bending tests and compression tests. A 30 mm × 20 mm × 20 mm wood block was cut from the bending sample with a significant difference in mechanical properties. Three to five pieces were cut from the block for the MFA test, and the rest were ground into powder for the determination of the relative crystallinity and crystallite size. Wood blocks with dimensions of 10 mm × 20 mm × 20 mm were extracted from bending samples with significantly different mechanical properties for anatomical and chemical analysis.

#### Mechanical properties testing

24 samples with dimensions of 20 mm (R) × 20 mm (T) × 300 mm (L) were used to test the MOR and MOE according to standard procedures [22, 23]. 24 samples with dimensions of 20 mm (R) × 20 mm (T) × 30 mm (L) were used to test the compressive strength parallel to the grain according to standard procedures [24]. Before testing, all the specimens were stored in a conditioning chamber (20 °C and 65% relative humidity) until they reached a moisture content of approximately 12%. The samples were oven-dried immediately after testing, and the precise moisture content was determined by comparing the weights measured before and after drying. All the mechanical property values were subsequently adjusted to 12% moisture content using the methodology described in the relevant standard procedures [22–24].
Anatomical analysis
To observe the anatomical structure, samples were transversely sectioned at a thickness of 15 µm using a sliding microtome, and the sections were dehydrated through an alcohol series, stained with safranine for photography and observed using a ZEISS Imager A1 microscope (ZEISS, Germany). At the same time, at least 50 tracheids were chosen to measure radial cell wall thicknesses by Axiovision image processing software (Axiovision Release 4.6.3, Carl Zeiss, Germany).

Crystallite size and relative crystallinity measurement
An X-ray diffractometer (X’Pert Pro, Panalytical, Netherlands) was used to measure the crystallite size and relative crystallinity. For testing, the powdery samples were placed in a sample box, flattened and placed horizontally in the instrument. Measurements were performed with CuKα radiation (λ = 1.541 Å, voltage = 40 kV, and current = 40 mA). The step size was 0.02°, and the scanning speed was 4°/min. The average width and length of the cellulose crystallites were calculated using Jade. The overlay of the measured curve and the fit curve were plotted, in addition to the estimated background and residual curve. Crystallite size estimation was done by using Scherrer’s equation after correcting for instrument resolution:

\[
B_{hkl} = \frac{K \lambda}{\cos \theta \sqrt{FWHM^2 - inst^2}},
\]

where \(B_{hkl}\) is the average size of the crystallites determined using hkl reflection, \(K\) is a constant (0.9), FWHM is the full width at half-maximum of the reflection, \(inst\) is instrumental broadening, and a Gaussian function presenting the reflection and amorphous background was fitted to the measured intensity. The average width and the average length of the crystallites were determined from the 002 and 040 reflections, respectively.

The relative crystallinity was determined by fitting the theoretical intensity of the crystalline cellulose via reflections presented by the Gaussian function and amorphous background to the measured intensity curve. The relative crystallinity index (CR) was determined as

\[
CR = \frac{I_{tot} - I_{amo}}{I_{tot}},
\]

where \(I_{tot}\) is the total experimental intensity and \(I_{amo}\) is the intensity of the amorphous background.

The relative crystallinity was determined for a few selected samples from intensities measured using both symmetrical transmission and reflection geometries. The relative crystallinity was calculated as a weighted sum of the crystallinity values so that a weight of 2/3 was used.

Fig. 1 Preparation of Pinus radiata samples. (A MOR and MOE. B Compressive strength parallel to the grain. C MFA, crystallite size, relative crystallinity. D Anatomical samples and chemical samples)
for the value determined from the intensity curve measured using symmetrical transmission and 1/3 for the value determined from the intensity curve measured using symmetrical reflection geometry [25, 26].

**MFA determination**

An X-ray diffractometer (X’Pert Pro, Panalytical, Netherlands) was used to measure the MFA of the wood chip samples. The sample was rotated, and the intensity curve was measured as a function of the rotation angle $\phi$ with a step of 0.5° and a measuring time of 180 s per point. A CuK$\alpha$ radiation source was used with $\lambda = 0.154$ nm, voltage $= 40$ kV, and current $= 40$ mA. The aperture of the incident beam was $2 \times 4$ mm. Mean MFA was determined according to the method developed by Cave [27], namely, 0.6 T, where T is an X-ray diffractometer parameter taken from the tangents drawn at the points of inflection [28, 29].

**Imaging FTIR spectroscopy**

To characterize the chemical differences in the cell walls of the tracheids, a Spectrum Spotlight 400 FTIR Imaging System (PerkinElmer Inc., Shelton, CT, USA) in attenuated total reflection mode was used. Slices approximately 14 μm thick were cut by a sliding microtome, and a Fourier transform infrared microimaging system was used to detect the in situ transmission mode. Each sample was measured at 2 locations of the corewood and outerwood, and 3 regions were randomly selected at each location. In the analysis of the infrared spectral data, at least 15 points were selected from each region to produce corresponding infrared spectral images for analysis. Six areas were measured from each of the outerwood and corewood locations.

**Results and discussion**

**Mechanical properties of outerwood and corewood**

**Bending properties**

The MOR and MOE are important indexes among the mechanical properties of wood. The MOR and MOE of *Pinus radiata* outerwood and corewood are listed in Table 1. By comparison, the MOR of the outerwood was 34.9% higher than that of the corewood. Statistical analysis showed that there was a significant difference in the MOR of the corewood and outerwood, and the average MOR was 92.8 MPa.

The average MOE was 9.26 GPa. Table 1 shows that the MOE of the *Pinus radiata* outerwood was 60.1% higher than that of the corewood. Statistical analysis showed that there was a significant difference in the MOE results between the corewood and outerwood.

**Compressive strength parallel to the grain**

As shown in Table 1, the outerwood had a higher compressive strength than those of the corewood. Statistical analysis showed that there was a significant difference in the compressive strength, and the average compressive strength of the corewood and outerwood was 34.0 MPa. The compressive strength parallel to the grain indicates the maximum capacity of wood to bear the pressure load along the grain direction, which is an important basis for the selection of components that are mainly subjected to pressure [29]. Therefore, the outerwood had a better compressive ability than the corewood.

The corewood and outerwood were in the vicinity of the pith and bark of *Pinus radiata*, respectively. The results showed that the mechanical properties of the MOR, MOE and the compressive strength parallel to the grain of the outerwood were obviously better than those of the corewood, which is similar to the findings of Alonso’s previous research [9], but contrary to the results observed for many hardwood species [30–32]. Attention should be given to the difference in mechanical properties between corewood and outerwood when processing and utilizing large-diameter *Pinus radiata*.

**The microstructures of outerwood and corewood**

**Anatomical characteristics**

Figure 2 shows the cross-sectional microstructures of the corewood and outerwood in *Pinus radiata*. The cell wall of the earlywood was large, and the cavity was ovoid, while the cell wall of the latewood was small and the

| Mechanics       | Number of samples | MOR (MPa) Mean | SD       | MOE (GPa) Mean | SD       | Compressive strength (MPa) Mean | SD   |
|-----------------|-------------------|---------------|---------|---------------|---------|---------------------------------|------|
| Outerwood       | 12                | 106           | 20.8    | 11.4          | 2.47    | 38.7                            | 7.48 |
| Corewood        | 12                | 78.9          | 9.69    | 7.12          | 1.36    | 29.3                            | 3.34 |
| *$P < 0.05$*    |                   |               |         | *             |         | *                               |      |

$*$P < 0.05 showed a significant difference
cavity was rectangular. The latewood cells of the outerwood were characterized by a small lumen and thick cell walls.

By measuring the radial double-wall thickness of tracheid cells of the corewood and outerwood of *Pinus radiata*, 3.65 ± 0.45 µm was obtained for the tracheid cells of the corewood, while 5.02 ± 2.83 µm was discovered for the tracheid cells of the outerwood. The results of variance analysis showed that there was a significant difference between corewood and outerwood. In previous study tracheids in ring 22 and ring 20 had significantly thicker cell walls than those in ring 7 of *Pinus radiata* [33]. The cell wall thickness at breast height of 19-year-old *Pinus radiata* rose by 57% from pith to bark [34] which agreed with our results in this study. The fiber wall thickness of poplar also increased gradually from the medulla to the bark [35]. The characteristics of low density (452 kg/m³) of corewood and high density (502 kg/m³) of outerwood in a previous study [5] were the same as the trend of cell wall thickness in this study.

**Relative crystallinity**

The diffraction patterns of the corewood and outerwood samples obtained by X-ray diffraction are shown in Fig. 3. Both the corewood and outerwood have three diffraction peaks near 18°, 22.5°, and 35° at 2θ, corresponding to the (101), (002) and (040) crystal planes, respectively. For (002), the maximum diffraction is at approximately 2θ = 22.5°, and the minimum value is at approximately 2θ = 18°. The X-ray diffraction patterns of the corewood and outerwood are basically the same, but the peak strength of the two clones at 2θ = 22.5° are significantly different, and the full widths at half-maximum are also different, indicating that the corewood and outerwood have the same cellulose crystal cells structure, while the degrees of crystallization are different. The values of the relative crystallinity of the outerwood and corewood are shown in Table 2. The results of the variance analysis showed that there was a significant difference in the crystallinity between the corewood and outerwood at the 0.05 level. The relative crystallinity of the outerwood was higher than that of the corewood.
Crystallite size
The peak segmentation fitting result of the X-ray diffraction data is shown in Fig. 4, and the curve has three characteristic peaks at $2\theta = 18^\circ$, 22.5°, and 35°, corresponding to the (101), (002) and (040) crystal planes, respectively.

As illustrated in Table 2, the mean crystallite width of the outerwood was larger than that of the corewood, the mean crystallite length of the outerwood was smaller than that of the corewood, while the differences in both the width and length were not significant. The mean crystallite width was 2.06 nm, slightly narrower than suggested in previous studies of 28-year-old *Pinus radiata*, 2.46 nm, from the denser outer part of the stem [36]. The crystallite sizes of Norway spruce were found to be different in different studies, probably because of the differences in the ages and locations of the samples [25, 28]. Similarly, no obvious change in the crystallite width from pith to bark was reported by Andersson [25].

Table 2 The relative crystallinity and crystallite size of *Pinus radiata* outerwood and corewood

| Sample location | Number of samples | Relative crystallinity (%) | Width (D002)/nm | Length (D040)/nm | Aspect ratio (D040/D002) |
|-----------------|-------------------|---------------------------|-----------------|-----------------|-------------------------|
| Outerwood       | 12                | 40.2 ± 0.07               | 2.14 ± 0.23     | 5.02 ± 12.5     | 2.36 ± 0.34             |
| Corewood        | 12                | 35.7 ± 0.04               | 1.98 ± 0.07     | 5.38 ± 15.1     | 2.73 ± 0.48             |

Microfibril angle
The X-ray diffraction patterns of samples from the corewood and outerwood are shown in Fig. 5. The measurement results showed that the mean MFA of the corewood was 22.3° ± 2.2° and that of the outerwood was 14.0° ± 1.54°. The results of the variance analysis showed that there was a significant difference between the corewood and outerwood at 0.05 level.

In studies of wood from different tree species, such as *Larix kaempferi* and white oak, the results showed that the mean MFA decreased gradually from the core to the bark along the radial direction [37, 38]. In Donaldson’s [39] study the MFA commonly showed larger values in the juvenile wood in the first few annual rings near the pith, followed by a decline to a more constant value in the mature wood. A study on the radial variation in MFA of *Pinus radiata* at 17 years old showed that the range of MFA from pith to bark was 30° to 11° [9]. The mean MFA of 8-month-old *Pinus radiata* was approximately 35° in Brennan’s study [40].

Chemical composition
In this study, infrared imaging was used to study the chemical composition of wood cell walls in situ, with an emphasis on the matrix materials. After obtaining the IR spectra of the cell wall, the average spectrum was baseline corrected at the absorption bands of 1800, 1548, 840, and 785 cm$^{-1}$ and normalized at 1424 cm$^{-1}$, corresponding to the CH$_2$ symmetric bending vibration of the CH$_2$-OH group in cellulose.

For lignin, the aromatic skeletal vibrations, together with the C=O stretch at 1598 cm$^{-1}$, the aromatic skeletal vibration at 1510 cm$^{-1}$, and the methoxyl C–H bending
band at 1366 cm⁻¹ were analyzed. Compared to the corewood, the outerwood showed an apparent increase in absorbance at 1510 cm⁻¹, while the absorbance at 1598, 1650 and 1366 cm⁻¹ was only marginally different (Fig. 6). The lignin types in the corewood and outerwood are the same. The increase in aromatic rings may be due to the higher lignin content in the outerwood. In Yang’s [41] study, the lignin structure of the 20-year-old Pinus radiata corewood was the same as that of the outerwood, and the lignin content of the outerwood was slightly higher than that of the corewood.

Correlation between microstructures and mechanical properties
In this study, the mechanical properties of the Pinus radiata corewood and outerwood were studied to provide basic data for their efficient utilization. Additionally, their microstructures, described by factors such as their anatomical parameters, relative crystallinity, crystallite size and MFA were explored. Some microstructures were significantly different, which provided a comprehensive description of the reason for the difference in the mechanical properties of Pinus radiata corewood and outerwood.

The relationship between anatomical parameters and mechanical properties
Wood is a material composed of a variety of cell types. The cell walls are the solid material of wood and the load-bearing structure of wood under stress. By comparing the microstructures of the Pinus radiata corewood and outerwood, it was found that the outerwood with a thicker cell wall, had a higher MOR, MOE and compressive strength parallel to the grain, while the corewood with a thinner cell wall had poorer mechanical properties. Tracheid wall thickness was positively correlated with the mechanical properties of Pinus radiata, indicating that the anatomical parameters may affect the mechanical properties at different radial locations in Pinus radiata.

The relationships among relative crystallinity, crystallite size, and mechanical properties
The growth process of trees is the process of the gradual accumulation of cellulose base fiber and an increase in crystallinity. By comparing the relative crystallinity and mechanical properties of corewood and outerwood, we found that with a higher relative crystallinity of the outerwood, the MOR, MOE and compressive strength parallel to the grain increased, whereas with a lower relative crystallinity of corewood, the MOR, MOE and compressive strength parallel to the grain decreased. Thus, the relative crystallinity could be a structural factor that affects the mechanical properties of Pinus radiata corewood and outerwood. As the relative crystallinity of cellulose increases, the binding forces between the molecular chains are strengthened and the orientation of the fibers becomes more favorable, which enhances the mechanical properties of most woods. Previous studies have shown that the degree of relative crystallinity is positively correlated with the dimensional stability and tensile strength of wood [9, 42].

The width and length of the crystallites were consistently positively related to the tangential and radial hardness, but negatively related to the transverse hardness in Damayanti’s [17] study. However, there was no significant difference in crystallite size found between the corewood and outerwood. Hence, the crystallite size is not the main factor affecting the bending properties and compressive strength parallel to the grain in this species of tree.

The relationship between MFA and mechanical properties
In this study, the mechanical properties MOR, MOE and compressive strength parallel to the grain of the Pinus radiata outerwood, were all higher than those of the corewood, while the MFA of the outerwood was lower than that of the corewood. This means that the MFA is negatively correlated with the mechanical properties, which is consistent with the conclusion of previous studies [33, 38, 43]. The arrangement of cellulose microfibrils in the cell walls plays a major role in determining the mechanical performance and shrinkage anisotropy of wood [44, 45]. The mechanical properties of outerwood are better than those of corewood in Pinus radiata. Thus, MFAs have an important influence on the mechanical properties of Pinus radiata.
The relationship between lignin and mechanical properties

Lignin is a high-molecular-weight aromatic polymer in the cell wall of trees and is one of the three components of the cell wall. Comparing the lignin of corewood and outerwood and the mechanical properties of wood, it was found that the outerwood had a slightly higher lignin content and a higher MOE, MOR, and compressive strength parallel to the grain. Previous studies have shown that lignin is packed into the cellulose skeleton and plays an important role in enhancing the mechanical strength of plants and maintaining the normal morphology of cells [46]. In a developing cell wall, cellulose fibrils coated with hemicellulose form an open network, whose empty spaces are progressively filled up with lignin [44, 47]. In this study, the lignin content may be an influencing factor of the mechanical properties of Pinus radiata, but not the major factor.

Conclusions

In this paper, a comparison was performed between the outerwood and corewood of Pinus radiata in terms of their mechanical properties and microstructures. The results showed that the MOR, MOE, and compressive strength parallel to the grain of the Pinus radiata outerwood were greater than those of the corewood, by 34.9, 60.1 and 32.3%, respectively. The relative crystallinity and the cell wall thickness of outerwood were 37.5 and 12.8% greater than those of the corewood, respectively. The MFA of the outerwood was 37.1% smaller than that of the corewood. FTIR spectroscopy images showed that the chemical composition and structure had no obvious change between the corewood and outerwood, but a slightly higher relative content of lignin was found in the cell walls of the outerwood. The crystallite size was not significantly different between the corewood and outerwood. The corewood and outerwood mechanical properties were positively correlated with the relative crystallinity and cell wall thickness, while, the MFA was negatively correlated with the mechanical properties. The synergy of these factors influence the mechanical properties of wood.

Abbreviations

MFA: Microfibril angle; MOR: Modulus of rupture; MOE: Modulus of elasticity; CR: Crystallinity index.

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