Thermodynamic analysis of binding of 4-methylumbelliferyl-α- and β-D-galactopyranosides to Momordica charantia lectin

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Binding of 4-methylumbelliferyl-α-D-galactopyranoside (MeUmbαGal) and the corresponding β-anomer (MeUmbβGal) to the Momordica charantia (bitter gourd) lectin (MCL) has been investigated by fluorescence spectroscopy. Binding of MeUmbαGal to MCL resulted in a decrease in the fluorescence intensity of the ligand. Saturation binding at 25°C resulted in a 17.8% decrease in the fluorescence intensity of the ligand. Quenching of the ligand fluorescence intensity was temperature-dependent and decreased with increase in temperature. Addition of lactose reversed the quenching due to binding, indicating that decrease in the fluorescence intensity of MeUmbαGal is due to the interaction of its carbohydrate moiety with the lectin. The changes in the fluorescence intensity of MeUmbαGal resulting from the binding were analysed to obtain the association constants for the process at different temperatures. At 25°C, the association constant, $K_a$, was determined to be $1.14 \times 10^4 \text{ M}^{-1}$. From the temperature dependence of the $K_a$ values the enthalpy and entropy of binding were estimated as $\Delta H^o = -25.9 \text{ kJ mol}^{-1}$ and $\Delta S^o = -9.1 \text{ J mol}^{-1} \text{ K}^{-1}$. A comparison of these values with the $\Delta H^o$ and $\Delta S^o$ values obtained for the binding of MeUmbβGal revealed that the higher affinity of the β-anomer is due to a larger enthalpy of binding, which overrides a larger negative entropy of binding for the latter.

LECTINS, the carbohydrate-binding proteins of non-immune origin, have been the subject of intense investigations in view of their interesting biological properties such as preferential agglutination of transformed cells, blood-group specificity, mitogenicity and hormone-mimicking activity. All these properties are manifested through the carbohydrate-binding activity of lectins, and hence it is of great interest to investigate their saccharide specificity and to delineate the forces that govern it. The Momordica charantia lectin (MCL) is a galactose-specific lectin present in the seeds of bitter gourd, which is used as a part of the diet in the tropics. MCL has been purified by affinity chromatography and characterized in considerable detail with regard to its macromolecular properties. Chemical modification studies have indicated that tryptophan and tyrosine residues are important for the sugar-binding activity of this lectin. Chemical modification, intrinsic fluorescence quenching and time-resolved fluorescence studies indicate that the tryptophan residues are in a heterogeneous environment, with at least two populations of tryptophan residues with different degrees of exposure being present in the protein. In addition to haemagglutination-inhibition studies, the binding of a few saccharides has been investigated by monitoring saccharide-induced changes in the intrinsic fluorescence emission spectrum of the protein. The binding of 4-methylumbelliferyl-β-D-galactopyranoside (MeUmbβGal) was investigated at different temperatures to monitor the changes in the fluorescence intensity of the ligand upon titration with MCL in order to determine the thermodynamic parameters associated with the binding, whereas the binding of 4-methylumbelliferyl-α-D-galactopyranoside (MeUmbαGal), the corresponding α-anomer was not studied. In this study the interaction of MeUmbαGal and MeUmbβGal with MCL was investigated at different temperatures, in order to delineate the thermodynamic basis for the higher affinity of the protein for the β-linked galactose moiety.

M. charantia lectin was purified by affinity chromatography on cross-linked guar gum. The protein gave a single band in PAGE and two bands corresponding to Mr ~28 kDa and ~30 kDa in SDS-PAGE, consistent with literature reports. Fluorescence measurements were performed on a Hitachi F-3010 fluorescence spectrometer, equipped with a water-jacketed cuvette holder that was maintained at constant temperature by means of a circulating water bath. Titration were performed by the addition of small aliquots of the protein from a concentrated stock solution (ca. 30 mg/ml) to 1.0 ml of a 5 μM solution of the fluorescent sugar. Samples were excited at 318 nm and emission spectra were recorded in the wavelength range of 330-450 nm. Slits of 5 nm were used for both excitation and emission monochromators. All the binding data reported here correspond to the average values obtained from two different titrations.

Fluorescence spectra of MeUmbαGal alone and in the presence of different concentrations of MCL, recorded at 25°C, are shown in Figure 1. From Figure 1 it is evident that addition of the lectin decreases the fluorescence intensity of MeUmbαGal. Further, changes in the fluorescence intensity resulting from protein addition could be reversed by the addition of lactose or galactose, demonstrating that the binding is mediated through the saccharide moiety of the labelled sugar. Similar results were for the titration of MeUmbβGal with MCL.

A binding curve, depicting the change in fluorescence intensity of the ligand (ΔF) as a function of the protein concentration, obtained from the above fluorescence spectra is shown in Figure 2. Here, it is seen that the binding curve displays saturation behaviour, clearly indicating that binding occurs at specific binding sites on the
protein. A plot of $1/\Delta F$ as a function of $1/[P]$, where $[P]$ is the total protein concentration, is given in the inset to Figure 2. The data yield a linear fit and from the $Y$-intercept of this plot, the change in fluorescence intensity at infinite protein concentration, $\Delta F_\infty$, was obtained. It was observed that the $\Delta F_\infty$ value was temperature-dependent and decreased with increase in temperature. At 15°C, the fluorescence intensity of MeUmbβGal decreased by 25%, whereas at 30°C, the decrease was only 15.2% (Table 1).

The titration data were then analysed according to eq. (1) in order to obtain the association constants, $K_a$ (ref. 11).

$$\log \frac{\Delta F/(F_c-F_\infty)}{[P]} = \log K_a + \log [P]$$

where $\Delta F$ is the fluorescence change at any point of the titration, $F_c$ is the fluorescence intensity at any point of the titration corrected for dilution, $\Delta F_\infty$ is the change in fluorescence intensity at infinite protein concentration and $[P]$ is the free protein concentration. A more detailed description of the method can be found elsewhere. Association constants for the binding of the umbelliferyl sugar ($K_a$) were obtained from the abscissa of the plot of $\log \{\Delta F/(F_c-F_\infty)\}$ vs $\log [P]$. A representative plot for the titration data obtained at 25°C is given in Figure 3. From the intercept of this plot the $K_a$ value has been estimated as $1.14 \times 10^4$ M$^{-1}$. Association constants for the MeUmbβGal-MCL interaction have been determined at several temperatures and the $K_a$ values obtained as described above are listed in Table 1.

As mentioned above, the binding of MeUmbβGal to MCL was investigated earlier by Khan et al. However, in order to compare the data obtained for both α and β-anomeric derivatives of the sugar under similar conditions and with the same lot of the purified protein, the binding of MeUmbβGal has also been investigated in this study. Experiments were performed essentially as described above for MeUmbαGal and it has been observed that the fluorescence intensity of the ligand is totally quenched at saturation binding at all temperatures investigated, an observation consistent with the results of Khan et al. The titration data were analysed as described above for the MeUmbβGal-MCL interaction, in order to determine the $K_a$ values for the binding of MeUmbβGal to MCL. At 25°C, the association constant was obtained as $2.12 \times 10^4$ M$^{-1}$, which is in excellent agreement with the value of $2.1 \times 10^4$ M$^{-1}$ obtained from equilibrium dialysis measurements, and also compares well with the value of $1.98 \times 10^4$ M$^{-1}$, obtained from fluorescence titrations by Khan et al. The $K_a$ values obtained at different temperatures are listed in Table 1.

![Figure 1](image1.png) **Figure 1.** Fluorescence spectra of MeUmbαGal in the absence and presence of *M. charantia* lectin at 25°C. Spectrum 1 is that of MeUmbαGal alone and spectra 2 to 13 were recorded after the addition of increasing concentrations of MCL.

![Figure 2](image2.png) **Figure 2.** Binding curve for titration of MeUmbβGal by MCL at 25°C. Change in fluorescence intensity at 376 nm (ΔF) was plotted as a function of the added protein concentration. (Inset) Plot of $1/(\Delta F)$ as a function of reciprocal protein concentration. From the $Y$-intercept of this plot, fluorescence intensity of the ligand at saturation binding was determined. See text for further detail.
From the $K_a$ values obtained at different temperatures it is clear that the interaction of MeUmbβGal with MCL is characterized by a stronger affinity than that of MeUmbαGal; the $K_a$ values for the former sugar being 1.6 to 2.0 times higher than the latter (Table 1). These results are consistent with the earlier observations that MeβGal is 1.5 times more potent than MeαGal in its ability to inhibit the haemagglutination by MCL$^9$.

The temperature dependence of the association constants for MeUmbαGal and MeUmbβGal has been analysed by Van’t Hoff plots of ln$K_a$ vs 1/T (see Figure 4) and linear fits have been obtained in each case. From the slope and intercept of these plots, enthalpy of binding ($\Delta H^\circ$) and entropy of binding ($\Delta S^\circ$) have been determined according to eq. (2):

$$\ln K_a = -\frac{\Delta H^\circ}{RT} + \frac{\Delta S^\circ}{R}.$$  

The enthalpy and entropy of binding for MeUmbαGal–MCL interaction were obtained as $\Delta H^\circ = -25.9$ kJ mol$^{-1}$ and $\Delta S^\circ = -9.1$ J mol$^{-1}$ K$^{-1}$, whereas the corresponding values for the binding of MeUmbβGal to MCL have been determined to be $\Delta H^\circ = -36.3$ kJ mol$^{-1}$ and $\Delta S^\circ = -39.3$ J mol$^{-1}$ K$^{-1}$. These values are also given in Table 1.

An analysis of the above thermodynamic parameters indicates that the enthalpy of binding for MeUmbβGal is considerably larger than the value obtained for MeUmbαGal, clearly indicating that binding of the β anomer is favoured by a larger enthalpy value. In addition, binding of the α anomer is associated with a smaller negative entropy compared to the β anomer. Therefore, the larger $\Delta H^\circ$ value associated with the binding of MeUmbβGal more than compensates for the negative

### Table 1. Maximum changes in fluorescence intensity ($\Delta F_\alpha$), association constants ($K_a$) and thermodynamic parameters ($\Delta H^\circ$, $\Delta S^\circ$) associated with the binding of MeUmbαGal and MeUmbβGal to *Monodora charantia* lectin, determined from the fluorescence spectral titrations

<table>
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<th>$T$ (°C)</th>
<th>$\Delta F_\alpha$ (%)</th>
<th>$K_a \times 10^3$ (M$^{-1}$)</th>
<th>$\Delta H^\circ$ (kJ mol$^{-1}$)</th>
<th>$\Delta S^\circ$ (J mol$^{-1}$ K$^{-1}$)</th>
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**Figure 3.** Chipman plot for binding of MeUmbαGal to MCL at 25°C. The Y-intercept of the plot yielded $pK_a$, from which the association constant, $K_a$ is determined.

**Figure 4.** Van’t Hoff plot for binding of MeUmbαGal (●) and MeUmbβGal (○) to MCL. Thermodynamic parameters, enthalpy of binding ($\Delta H^\circ$) and entropy of binding ($\Delta S^\circ$) were determined from the slope and intercept respectively, of this plot according to eq. (2). See text for further detail.
Bud break and plantlet regeneration in vitro from mature trees of Pinus roxburghii Sarg.

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Terminal and axillary buds of 30-year-old mature Pinus roxburghii Sarg. trees were collected to test their in vitro response. Five different types of shoots containing these buds were available in a year. Shoots collected between February and May gave the best response. Bud sprouting could be obtained on solid half strength DCR basal medium (1/2 DCR), fortified with BA from 0.44 to 11.1 μm, 11.1 μm being the optimum concentration used. On taking these buds for elongation, however, only those transferred from 0.44 μm elongated on plain 1/2 DCR after three passages of 45 days each. The initial concentration of BA therefore had an influence on the elongation of shoots. Rooting was obtained in 5% of the shoots on the elongation medium. Optimum rooting (54%) was obtained after a 24 h treatment on 12.25 μm IBA and the plantlets survived upon transfer to polyhouse.

PINUS roxburghii Sarg., commonly known as ‘chir pine’, is one of the five species of pines occurring wild in the Himalaya and hills of Assam in India† and in the monsoon belt of northeastern Pakistan to Bhutan. It grows to a height of 45 to 54 m and is known to be fairly drought-resistant. The wood is widely used for making furniture, door and window frames, railway sleepers, paper, pulp, etc. Oleoresin from chir pine is the main source of turpentine in India, which is chiefly used as a solvent for thinning, paints and varnishes besides having medicinal uses†. Natural regeneration is through seeds. Trees less than 30 years old rarely bear cones. A good seed year occurs only once in four to five years. Considering its uses, seeding problems and depletion of the natural stocks, there is an urgent need to have alternative methods for the establishment of plantation. This can be achieved through two methods of vegetative propagation. One involves rooting of juvenile cuttings from seedlings raised from hybrid seeds, as is done in the case of radiata pine. The other involves rooting of stem cuttings from superior phenotypes. Application of tissue culture techniques would be another method for large-scale cloning of mature trees. In the case of conifers this is more important, as the rooting frequency declines when the parent plant is more than ten years old.

In the last 15 years, there have been several reports on in vitro propagation of conifers. In recent years, there has been success in conifer propagation through both somatic embryogenesis 6–9 and induction of adventitious buds10–13. Most of these reports deal with studies using juvenile explants (seeds, seedlings, young cotyledons, etc.). However, there are few reports on the in vitro propagation from mature tree-derived explants14–16. The large genome sizes of pines and the recalcitrance of tissues from mature trees to in vitro manipulation continue to present challenges to researchers17. In the case of P. roxburghii, only adventitious-bud proliferation has been reported so far18. To our knowledge, there are no reports on in vitro vegetative propagation of this species from mature tissues.

The present study is part of an ongoing programme to develop micropropagation methods for Pinus species.


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