Research Paper

Calculated and Experimental Infrared Spectra of Substituted Naphthoquinones

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Abstract

In recent years there has been interest in incorporating high-potential, halogenated 1,4-naphthoquinones (NQs) into the A\(_1\) binding site in photosystem I (PSI) photosynthetic reaction centers. This interest in part stems from the considerably altered bioenergetics of electron transfer that occur in PSI with such a substitution. FTIR studies of PSI complexes with halogenated NQs incorporated are also being undertaken. With this in mind it is necessary to consider FTIR absorption spectra of these halogenated quinones in solution, and here we present FTIR absorbance spectra for 2-chloro-1,4-naphthoquinone (2ClNQ), 2-bromo-1,4-naphthoquinone (2BrNQ), 2,3-dichloro-1,4-naphthoquinone (Cl\(_2\)NQ) and 2,3-dibromo-1,4-naphthoquinone (Br\(_2\)NQ) in tetrahydrofuran (THF). The FTIR spectra of the substituted naphthoquinones (NQs) were compared to FTIR spectra of 2-methyl-3-phytyl-1,4-naphthoquinone (phyloquinone (PhQ)) and 2-methyl-naphthoquinone (2MNQ). To aid in the assignment of bands in the experimental spectra, density functional theory (DFT) based vibrational frequency calculations for all the substituted NQs were undertaken. The calculated and experiment spectra agree well. By calculating normal mode potential energy distributions unambiguous, quantitative band assignments were made. The calculated and experimental spectra make several predictions about what might be observed in time resolved FTIR difference spectra obtained using PSI with the different quinones incorporated.

Keywords

Naphthoquinone, Phylloquinone, FTIR, Density functional theory, vibrational frequency calculations

Abbreviations: FTIR, Fourier transform infrared; H-bond, hydrogen bond; NQ, 1,4-naphthoquinone; PSI, photosystem I; PhQ, phylloquinone (2-methyl-3-phytyl-1,4-naphthoquinone); PED, potential energy distribution; 2ClNQ, 2-chloro-1,4-naphthoquinone; 2BrNQ, 2-bromo-1,4-naphthoquinone; Cl\(_2\)NQ, 2,3-dichloro-1,4-naphthoquinone; Br\(_2\)NQ, 2,3-dibromo-1,4-naphthoquinone; THF, tetrahydrofuran; DFT, density functional theory.
**Introduction**

The redox properties of quinones allow them to be utilized in a wide range of functions in different biological systems. In photosynthetic reaction centers (RCs), quinones are well known cofactors in electron transport (ET). In type II RCs, which includes photosystem II (PSII) of oxygen evolving organisms and purple, nonsulfur bacteria, quinones function as terminal electron acceptors in the so-called QA and QB binding sites. Even although the QA and QB binding sites may be occupied by the same quinone (such as plastoquinone in PSII their functions are quite different, where the QA quinone is an intermediary in ET while the QB quinone is involved in proton-coupled ET. In type I RCs, which have iron sulfur clusters as terminal electron acceptors, and includes photosystem I (PSI) from plants and algae, quinones function as an intermediary in transferring electrons from A0 to Fx.

In native PSI a phylloquinone (PhQ) molecule occupies the A1 binding site. PhQ is a di-substituted 1,4-naphthoquinone (NQ) (2-methyl,3-phenyl-NQ) (also known as Vitamin K1). Menaquinones (MQs) are identical to PhQ except in the degree of saturation of the hydrocarbon tail, and MQs are found in the QA binding site in type II purple bacterial reaction centers from Blastochloris viridis and Chloroflexus aurantiacus. The can also be substituted into the QA binding site in purple bacterial reaction centers from Rhodobacter sphaeroides. The function of substituted NQs in both type I and II RCs is therefore of some interest. In particular, exactly how the substituent groups of the NQs can alter or modulate their redox properties of quinones in protein binding sites is of considerable interest.

In recent years it has become possible to substitute a range of different NQs into the A1 binding site in PSI. These PSI particles with different NQs incorporated into the A1 binding site are now being studied using FTIR difference spectroscopy, and of particular interest for the purposes of this manuscript is PSI with different halogenated NQs incorporated. The overall goal in FTIR studies of PSI with different NQs incorporated is the identification of bands in the spectra that are associated with the neutral and reduced states of the incorporated NQs. These bands will provide information on pigment protein interactions, that may help modulate the redox properties of the incorporated quinones.

Study of halogenated NQs in the A1 binding site is very interesting, not only in terms of the FTIR difference spectra, but also in terms of the bioenergetics, which are very different. Halogenated NQs intrinsically have a very high redox potential, and even in the A1 binding site in PSI, the P700A1− radical pair state is considerably lower in energy than the P700F− state, making uphill ET from A1− to Fx essentially impossible. Therefore, the P700A1− state decays by radical pair recombination, which occurs on a tens to hundreds of microsecond timescale at both 298 and 77 K. This simplifies the collection of highly-sensitive time resolved (TR) FTIR DS.

When considering bands in TR FTIR DS obtained using PSI with different NQs incorporated, the starting point is to first obtain and interpret the FTIR spectrum of the NQ in solution. Such an interpretation invariably involves (these days) the use of quantum chemical computational methods to simulate the experimental spectra. These suggested initial steps are the subject of this manuscript.

In this manuscript FTIR absorbance spectra for 2-chloro-1,4-naphthoquinone (2ClNQ), 2-bromo-1,4-naphthoquinone (2BrNQ), 2,3-dichloro-1,4-naphthoquinone (Cl2NQ), and 2,3-dibromo-1,4-naphthoquinone (Br2NQ) in tetrahydrofuran (THF) are obtained. These spectra are compared to that obtained for PhQ (and DMNQ) and 2MNQ in THF. In addition, in order to help assign bands in the experimental spectra density functional theory (DFT) based vibrational frequency calculations are undertaken.

**Materials and methods**

All quinones and solvents used here were obtained from Sigma-Aldrich Inc. (St Louis, MO), and used as received. 2ClNQ was obtained from Pure-Chemistry Scientific (Burlington, MA) and used as received. FTIR absorption spectra of the NQs were measured using a Varian 600 UMA FTIR
microscope (Varian Inc., MA, USA), utilizing a mercury cadmium telluride detector. Spectra were collected at 4 cm$^{-1}$ resolution. The different NQs were dissolved in tetrahydrofuran (THF) at various concentrations, and a small drop (<5 μL) was deposited on a zinc selenide window. An FTIR spectrum of pure THF was obtained and interactively subtracted.

Calculations were undertaken using Gaussian16 software.\textsuperscript{18} (Gaussian Inc. Wallingford, CT). The phytyl tail of PhQ was truncated to a 5-carbon unit [CH$_2$CHC(CH$_2$)$_2$]. DFT calculations were performed using the B3LYP functional in combination with the 6-31+G(d) basis set. The solvent was considered using the integral equation formalism (IEF) of the polarizable continuum model (PCM),\textsuperscript{19} as it is implemented in Gaussian16. Harmonic, normal mode vibrational frequency calculations result in “stick spectra”, which we convolve with Gaussian functions of 4 cm$^{-1}$ half-width, to produce more realistic looking calculated spectra.

Assignment of calculated vibrational frequencies to molecular groups is often based on the calculation of potential energy distributions (PEDs) associated with the normal modes. PEDs are calculated using VEDA.\textsuperscript{20} PEDs provide an estimate of how much a specific molecular group vibration contributes to a normal mode.

**Results and Discussion**

Figure 1 shows the structure and numbering for the six quinones considered in this study. PhQ is the same as 2MNQ except that the phytyl chain is replaced with a hydrogen atom. Figure 2 shows FTIR absorption spectra in the 1750-1200 cm$^{-1}$ spectral region for PhQ, 2MNQ, 2CINQ, 2BrNQ, Cl$_3$NQ, and Br$_2$NQ in THF. A spectrum for PhQ cast in a film of hexane has been obtained previously, and is very similar to that presented in Figure 2.\textsuperscript{21} A spectrum for 2,3-dimethyl-1,4-naphthaquinone (DMNQ) in CCl$_4$ has also been obtained.\textsuperscript{22} DMNQ is the same as PhQ except the phytyl chain has been replaced with a methyl group, and it is found that the FTIR absorption spectra of PhQ and DMNQ (in the 1680-1580 cm$^{-1}$ region) are nearly identical.

Figure 3 shows DFT calculated FTIR spectra for the six quinones considered in this study. The PEDs associated with the normal modes that give rise to the most intense bands in the spectra in Figure 3 are listed in Table 1. The calculated and experimental spectra clearly display similarities in both peak positions and in relative band intensities. How the calculated and experimental bands are associated is indicated in Table 1. In this manuscript calculated frequencies are scaled by 0.978, which is arbitrarily chosen so that the observed band at 1662 cm$^{-1}$ for neutral PhQ matches the frequency for the calculated band.

**Modes of the phytyl chain of PhQ.** PhQ in THF (Figure 2A) displays bands at 1469 and 1377 cm$^{-1}$ that are absent in the spectra of the other NQs (Figure 2). These bands are therefore likely to be associated with the hydrocarbon phytyl chain of PhQ. The calculations support this notion, and show that the 1469 cm$^{-1}$ band is due to CH$_2$ bending modes [$\delta$(CH$_2$)] of the phytyl-tail, while the 1377 cm$^{-1}$ band is due to both $\delta$(CH$_2$) and $\delta$(CH$_3$) modes of the phytyl-tail (Table 1). This conclusion also agrees with previous studies.\textsuperscript{23,24} These assignments indicate the appropriateness of the computational methods used here.

**Quinone C=O vibrations.** For PhQ in THF the most intense absorption band is found at 1662 cm$^{-1}$, and calculations indicate that this band is due to the antisymmetric stretching vibration of both C=O groups (C$_2$=O, 54% and C$_3$=O, 30%) (Table 1). For 2MNQ in THF a similar intense peak is observed at 1666 cm$^{-1}$ with a weak shoulder near 1670 cm$^{-1}$ (Figure 2B). Computationally the intense 1666 cm$^{-1}$ band is also assigned to an antisymmetric stretching vibration of both C=O groups (C$_1$=O, 52% and C$_3$=O, 32%) (Table 1). Similarly, for Cl$_2$NQ and Br$_2$NQ in THF intense bands are observed at 1686 and 1684 cm$^{-1}$, respectively. This is an upshift of 26 and 22 cm$^{-1}$ relative to the same band of PhQ. Importantly, the observed upshift is very similar to that predicted computationally (26 and 24 cm$^{-1}$). In addition, the normal mode giving rise to the intense bands at 1686 and 1684 cm$^{-1}$ for Cl$_2$NQ and Br$_2$NQ is again calculated to be an antisymmetric coupled vibration of both the C$_1$=O and C$_3$=O groups (For Cl$_2$NQ: C$_1$=O, 44% and C$_3$=O, 44%. For Br$_2$NQ: C$_1$=O, 44% and C$_3$=O, 45%) (Table 1).
Figure 1: Structure and numbering of the different NQs considered here.

Figure 2: FTIR absorption spectra for (A) PhQ, (B) 2MNQ, (C) 2CINQ, (D) 2BrNQ, (E) Cl₂NQ, and (F) Br₂NQ in THF. The spectra were scaled (1695-1640 cm⁻¹) so that the intensities of the bands around 1684 cm⁻¹ are similar.
Figure 3: Calculated IR absorption spectra for (A) PhQ, (B) 2MNQ, (C) 2CINQ, (D) 2BrNQ, (E) Cl₂NQ, and (F) Br₂NQ in THF. Frequencies are scaled by 0.978. Spectra were scaled so that the intensities of the bands between 1688-1662 cm⁻¹ are similar. The actual intensities (in km/mol) are indicated in Table 1.

Table 1: Calculated normal mode vibrational frequencies (in cm⁻¹) and intensities (in km/mol) for the six NQs in THF. Only the most intense normal modes are listed. Frequencies are scaled by 0.978 to compare to experimental band frequencies, which are also listed. Potential energy distributions (PED) (in %) are also listed. Only molecular groups with PED above 10% are included. Abbreviations: ν, bond stretching; δ, bending vibration; a, aromatic part of NQ ring; q, quinonic part of NQ ring; H*, hydrogen atom attached at position C₃. AS, antisymmetric.

<table>
<thead>
<tr>
<th>Experimental frequencies (in THF) (cm⁻¹)</th>
<th>Calculated frequencies (in THF) (cm⁻¹)</th>
<th>Intensity (km/mol)</th>
<th>Potential Energy Distribution (%)</th>
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<td><strong>PhQ</strong></td>
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<tr>
<td>1662</td>
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<td>1597</td>
<td>1597</td>
<td>204.7</td>
<td>ν(C-C)ₐ 64%</td>
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<td>1469</td>
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<td>114.6</td>
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<td>1597</td>
<td>210.6</td>
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<td>ν(C-C)ₐ 16% + δ(CCH)ₐ 22%</td>
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For 2CINQ and 2BrNQ in THF the main intense peak discussed above is split into two separate peaks at 1683 and 1668 cm⁻¹ (Figure 2C, D). Such a clear (15 cm⁻¹) splitting is in agreement with calculations, which indicate intense bands at 1688-1690 and 1668-1669 cm⁻¹ (Figure 3C, D). The 1683 and 1668 cm⁻¹ bands for 2CINQ and 2BrNQ in THF are predicted to be due to C₁=O (86%) and C₄=O (83%) vibrations (Table 1). This is an interesting result, as it is not observed or calculated for 2MNQ, which one might anticipate to have spectra more similar to that of 2CINQ or 2BrNQ than that for Cl₂NQ or Br₂NQ.

### Quinonic and aromatic ring C=C vibrations.
For PhQ in THF an absorption band is observed at 1619 cm⁻¹ (Figure 2A). A similar band is calculated at 1624 cm⁻¹ (Figure 3A), and is predicted to be mainly due to a C₂=C₃ stretching vibration (ν(C₂=C₃), 68%) (Table 1). For 2MNQ in THF the corresponding band is upshifted 6 cm⁻¹ compared to PhQ to 1625 cm⁻¹ (ν(C₂=C₃), 68%) (Table 1). However, the calculated upshift is 15 cm⁻¹ (Table 1).

For PhQ in THF an absorption band is observed at 1597 cm⁻¹ (Figure 2A). A similar band is calculated at the same frequency (Figure 3A), and is predicted to be mainly due to a C=C stretching associated with the aromatic part of the NQ ring (ν(C=C)ₙ, 64%) (Table 1). For 2MNQ in THF the corresponding band is downshifted 2 cm⁻¹ compared to PhQ at 1595 cm⁻¹ (ν(C=C)ₙ, 65%) (Table 1). This agrees with calculations which predict the ν(C=C)ₙ mode to be at the same frequency for both 2MNQ and PhQ (Table 1).

For PhQ and 2MNQ the ν(C₂=C₃) vibration is at a higher frequency than the ν(C=C)ₙ vibration. This appears to be true for 2BrNQ and 2CINQ, although the reverse is calculated to be true for Cl₂NQ and Br₂NQ:

### Table 1

<table>
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<tr>
<th></th>
<th>2CINQ</th>
<th></th>
<th>2BrNQ</th>
<th></th>
<th>Cl₂NQ</th>
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<td>ν(C₁=O)</td>
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<td>ν(C₁=O)</td>
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<td>ν(C₁=O)</td>
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vibration. The corresponding bands for Cl$_2$NQ and Br$_2$NQ in THF (at least from a visual inspection of the spectra) are observed at 1593 and 1591 cm$^{-1}$, respectively. Corresponding bands appear to be calculated at very similar frequencies (1595 and 1596 cm$^{-1}$). These bands are due to the $\nu$(C=C)$_q$ vibration, however.

For 2CINQ and 2BrNQ in THF bands are observed at 1578 and 1575 cm$^{-1}$, respectively (Figure 2). Corresponding bands appear to be calculated at 1596 and 1597 cm$^{-1}$. These bands are due to the $\nu$(C=C)$_a$ vibration. The corresponding bands for Cl$_2$NQ and Br$_2$NQ in THF (at least from a visual inspection of the spectra) are observed at 1566 and 1556 cm$^{-1}$, respectively. Corresponding bands appear to be calculated at very similar frequencies (1567 and 1559 cm$^{-1}$). These bands are due to the $\nu$(C$_2$=C$_3$) vibration, however.

In summary, all of the experimental and calculated spectra show three or four bands in the 1550-1690 cm$^{-1}$ region. The highest frequency bands are due to C$_1$=O and C$_6$=O vibrations, or an antisymmetric combination of both. The next two lower intensity bands are due to $\nu$(C$_2$=C$_3$) and $\nu$(C=C)$_a$ vibrations, with the $\nu$(C$_2$=C$_3$) vibration being higher in frequency than the $\nu$(C=C)$_a$ vibration for PhQ, 2MNQ, 2CINQ and 2BrNQ, and lower for Cl$_2$NQ and Br$_2$NQ. One noteworthy point is that for Cl$_2$NQ in THF the FTIR spectrum in Figure 2E displays shoulders on the high frequency sides of some of the above discussed absorption bands. These shoulders are of unknown origin at present. A second noteworthy point is that the relative intensities of the three (or four) bands in the experimental spectra overall agree reasonably well with the calculated spectra.

PhQ in THF displays an intense band at 1297 cm$^{-1}$, which is at least as intense as the 1662 cm$^{-1}$ band (Figure 2A). Such an intense band is calculated at 1280 cm$^{-1}$, and is predicted to be due to $\nu$(C-C)$_a$ (10%) and $\nu$(C-C)$_q$ stretching (39%). Most TR FTIR DS experiments of native PSII have focused on a spectral region above 1400 cm$^{-1}$. However, the calculated and experimental spectra presented here indicate that extending work to lower frequency (to ~ 1200 cm$^{-1}$) might be useful for the identification of a band of neutral PhQ in the A$_1$ binding site. Prior to undertaking this experiment, however, it will off course be useful to calculate spectra for PhQ$^-$ to establish how and if this band is altered upon anion formation.

The PhQ band calculated at 1280 cm$^{-1}$ in Figure 3A is found downshifted ~15 cm$^{-1}$ to 1257-1254 cm$^{-1}$ for Cl$_2$NQ and Br$_2$NQ (Figure 3E, F). This calculated result suggests the bands of 1277 and 1267 cm$^{-1}$ in the Cl$_2$NQ and Br$_2$NQ FTIR spectra in Figure 2E and F correspond to the 1297 cm$^{-1}$ band for PhQ. This result, suggesting an intense band of neutral PhQ near 1295 cm$^{-1}$, and corresponding bands of neutral Cl$_2$NQ and Br$_2$NQ -~20-30 cm$^{-1}$ lower in frequency, may be extremely useful and help in the identification of bands in future TR FTIR DS obtained using PSI with neutral Cl$_2$NQ and Br$_2$NQ incorporated. The value of this result is even greater when one considers that in the past it has proven very difficult to unambiguously identify absorption bands associated with the neutral quinones incorporated into PSI.

It appears that the calculated band of PhQ at 1280 cm$^{-1}$ is split into two bands for 2CINQ and 2BrNQ. Such a splitting is also found in the 2MNQ calculated spectra. For 2CINQ and 2BrNQ bands are found at 1290-1288 and 1248 cm$^{-1}$. The former band is upshifted ~9 cm$^{-1}$ (relative to the 1280 cm$^{-1}$ band of PhQ) and is due to $[\nu$(C-C)$_q$ (32%) + $\delta$(CCH)$_q$ (12%) + $\delta$(CCH)$_a$ (14%)]. The latter band is downshifted 32 cm$^{-1}$ and is due to $[\nu$(C-C)$_a$ (10%) + $\delta$(CCH)$_a$ (25%)]. These calculated spectra suggest that the bands at 1290-1288 and 1248 cm$^{-1}$ could correspond to the bands at 1297-1294 and 1252-1248 cm$^{-1}$ in the experimental spectra for 2CINQ and 2BrNQ in THF. Again, the calculated and experimental spectra presented here predict that bands of neutral 2BrNQ and 2CINQ may be found at ~1295 and ~1250 cm$^{-1}$ in TR FTIR DS obtained using PSI with 2BrNQ and/or 2CINQ incorporated into the A$_1$ binding site. We will certainly be on the lookout for such features in the TR FTIR DS when they become available.

Conclusions

FTIR absorption spectra for six NQs in THF are presented. Corresponding calculated spectra are also presented, and indicate detailed normal mode assignments for the bands in the
experimental spectra. For PhQ in THF a band is observed at 1469 cm\(^{-1}\), which is absent in FTIR spectra for the other quinones, indicating that this band is due to vibrations associated with the phytyl tail of PhQ.

Neutral PhQ/2MNQ/Cl\(_2\)NQ/Br\(_2\)NQ in THF gives rise to an intense absorption band at 1662/1666/1686/1684 cm\(^{-1}\), respectively, that calculations indicate is due to the antisymmetric coupled vibration of both C=O groups. For 2CINQ/2BrNQ in THF this band is split into two bands, the higher/lower frequency band being due to the C1=O/C3=O stretching vibration, respectively.

FTIR spectra of PhQ/Cl\(_2\)NQ/Br\(_2\)NQ in THF display an intense absorption band at 1297/1277/1267 cm\(^{-1}\), respectively, that calculations indicate is due to a C-C stretching of both the aromatic and quinonic parts of the NQ ring. For 2MNQ/2CINQ/2BrNQ in THF this band is split into two. The spectra presented here suggest or lead to predictions about the location of quinone bands that may be found in TR FTIR DS.

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