A double co-sensitization strategy using heteroleptic transition metal ferrocenyl dithiocarbamate phenanthroline-dione for enhancing the performance of N719-based DSSCs†

Amita Singh, Devyani Srivastava, Suresh W. Gosavi, Ratna Chauhan,* Muthupandian Ashokkumar, Awad Naseer Albalwi, Mohd. Muddassir and Abhinav Kumar

Three new heteroleptic dithiocarbamate complexes with formula [M(Phen-dione)(Fcdtc)]PF₆ (where M = Ni(II) Ni-Fc, Cu(II) Cu-Fc) and [Co(Phen-dione)(Fcdtc)₂]PF₆ (Co-Fc) (Fcdtc = N-ethanol-N-methyl(ferrocene dithiocarbamate and Phen-dione = 1,10-phenanthroline-5,6-dione; PF₆⁻ = hexafluorophosphate) were synthesized and characterized using microanalysis, FTIR, electronic absorption spectroscopy and mass spectrometry. The solution state electronic absorption spectroscopy for all three complexes displayed a band at ~430 nm corresponding to the ferrocene unit and another low-intensity band in the visible region arising because of the d–d transitions. These newly synthesized complexes were used as co-sensitizers for the state-of-the-art di-tetrabutylammonium cis-bis(isothiocyanato)bis(2,2′-bipyridyl-4,4′-dicarboxylato)ruthenium(II) (N719) dye in dye-sensitized solar cells (DSSCs). Among the three co-sensitizers/co-adsorbent-based DSSC set-ups, the assembly fabricated using Co-Fc/N719 displayed good photovoltaic performance with 5.31% efficiency (η) while a new triple component strategy inculcating N719, Co-Fc and Cu-Fc dyes offered the best photovoltaic performance with 6.05% efficiency (η) with incident photon to current conversion efficiency (IPCE) of 63%. This indicated an upliftment of the DSSC performance by ~38% in comparison to the set-up constructed by employing only N719 dye (η = 4.39%) under similar experimental conditions.

Introduction

There is a rising concern about fulfilling the energy needs of the exponentially increasing world population.¹ Hence, apart from conventional non-renewable sources of energy, scientists and industries working in the energy sector are looking for renewable sources of energy. Amongst these, solar energy is a prudent option that can fulfill the energy demand of the entire world as annually around 6000 times more energy reaches from the sun to the earth’s surface than the energy consumed by the entire world.² To convert solar radiation into accessible form, apt photovoltaic devices that can capture and convert the quanta coming from the sun to electrical energy for their concomitant utilization in a plethora of applications are required. However, the photovoltaic devices developed using solid-state p–n junctions are expensive and difficulties associated with the fabrication of the p–n junction are the main obstacles in their use. Hence, due to potentially low production costs and good efficiency, dye-sensitized solar cells (DSSCs) have received considerable attention from researchers working in this field.³ Hence, after the pioneering work by Grätzel et al. in 1991, during the past twenty years, DSSCs had become the alluring class of devices due to their ease of fabrication with excellent incident photon conversion efficiency, charge transfer flexibility, transparency, and colour properties.⁴–⁷ A typical DSSC set-up is designed using a photo-anode fabricated using an appropriate light harvester coated over a suitable semiconducting material, especially nanocrystalline TiO₂, a counter electrode, and a redox mediator.⁸

The most important component of any typical DSSC set-up is the photosensitizer and amongst which the Ru(n) polyppyridyl sensitizer, especially state-of-the-art di-tetrabutylammonium
cis-bis-(isothiocyanato)-bis-(2,2′-bipyridyl-4,4′-dicarboxylato)-ruthenium(II) (N719) dye exhibited certified photovoltaic efficiency of 11%.\(^7\) In spite of its low abundance, high cost, poor absorption in the near-infrared (NIR) region, and difficulty in purification and isolation of Ru(II) complexes, scientists working in the area of photovoltaics continuously developed analogues of N719.\(^8\) Apart from ruthenium-based sensitizers, till date, numerous organic and inorganic photosensitizers have been reported wherein the efficiency with metal-free organic sensitizers reached 14%.\(^9\)–\(^16\) Also, to uplift the photovoltaic performance, porphyrin-based sensitizers have been used due to their high molar extinction coefficients.\(^17\)–\(^18\) Also, by judicious modification in the structure of heterocycles such as pyrroles, thiophenes, imidazoles, their electrochemical and optical properties can be tuned. This results in broad electronic absorption, which ranges from the visible to the NIR region.\(^19\)–\(^26\) The pertinent examples include meso-phenyl carboxylate zinc(II) porphyrin and porphyrin substituted with carboxylic groups at the β-pyrrole position that offered overall efficiencies of 3.5% (ref. 21) and 7.1%, respectively.\(^23\) Also, recently Grätzel \textit{et al.} delineated YD2-O-C832 and SM315 sensitizers that offered efficiencies of up to 12.5% and 13%, respectively.\(^27\)–\(^28\) Additionally, D–π–A porphyrin-based sensitizers with phenothiazine as an electron donor and benzothiadiazole as an acceptor exhibited ~10% photovoltaic efficiency.\(^29\) While, porphyrin-based organic co-sensitizers along with N719 dye exhibited 14.2% photovoltaic efficiency.\(^30\)–\(^31\) Giribabu \textit{et al.} reported D–π–A based porphyrin-based sensitizers \textit{viz.} LG24, LG26, LG25, and LG27 exhibit ~10.45% power conversion efficiency.\(^30\)–\(^31\) Apart from these reports, the use of 2D and 3D hybrid systems \textit{viz.} perovskite solar cells constitute very promising photovoltaic devices with a maximum overall conversion efficiency of 18.8%.\(^32\) Nevertheless, their instability and higher charge recombination hampered their commercial utilization. Further, photosensitizers having heterocyclic units \textit{viz.} pyranylidene, thienothiophenes, carbazole, oxadiazole, benzimidazole and dicyanovinyl, dicyanoimidazole, tricyanofuran, indandione, (thio)barbituric acid have been widely employed as push–pull units in DSSCs.\(^33\)–\(^34\) In relation to this, metalloccenes, ferrocene (Fc), and Fc-based systems acting like electron-rich moieties can be employed as sensitizers in DSSC setups.\(^34\) Hence, a range of coordination complexes and compounds comprising ferrocene entities have been used as sensitizers in DSSCs.\(^35\) Further, to extend the applicability of ferrocene as a photosensitizer in DSSC and to uplift its stability and performance, π-conjugated ferrocene and ferrocene-appended heterocycles have been reported.\(^35\)

Apart from these reports, the ferrocene-based complexes have been utilized as co-sensitizers to enhance the DSSC performance of the state-of-the-art N719 dyes.\(^36\) These investigations laid a firm platform to utilise ferrocene-based derivatives as co-sensitizers in DSSCs. The 1,10-phenanthroline-5,6-dione ligand-based complexes display the potential to exhibit higher stability and have the capability to allow control over redox tenability and also these can engender suitable metal-loligands for the generation of functionalized coordination complexes that could be utilized as sensitizers and co-sensitizers in DSSCs.\(^37\)

In view of the aforementioned aspects associated with the ferrocene functionalized coordination complexes and in our quest to develop new ferrocene functionalized sensitizers, in the presented report new-heteroleptic complexes inculcating ferrocene and 1,10-phenanthroline-5,6-dione have been synthesized and their plausible applications as sensitizers and co-sensitizers have been delineated. The outcomes of these investigations are presented hereafter.

### Results and discussion

#### Syntheses

The ferrocene appended dithiocarbamate ligand was obtained by condensation of ferrocene-carboxaldehyde and ethanolamine that yielded a Schiff base, which after reduction with sodium borohydride gave ferrocene functionalized secondary amine. This reaction with KOH and carbon disulfide yielded the potassium salt of dithiocarbamate ligand (Scheme 1). Finally, reacting a methanol solution of [M(Phen-dione)] (M = Co(II), Ni(II), and Cu(II)) with the dithiocarbamate ligand yielded the desired sensitizers Co-Fc, Ni-Fc, Cu-Fc (Scheme 1). All three sensitizers were air and moisture stable and were characterized by microanalysis, FTIR, UV-Vis, and mass spectrometry.

The molecular structure of all three sensitizers was assessed with the help of theoretical calculations and optimized geometries are presented in Fig. 1. The calculations revealed that the immediate geometry around Ni(II) in Ni-Fc and Cu(II) in Cu-Fc are distorted square planars, whereas in Co-Fc the geometry around Co(II) is distorted octahedral (Fig. 1). The geometry

---

**Scheme 1** Synthetic route for the co-sensitizers.
around the Ni([n] and Cu([n] are defined by two sulfur centers of Fe-dtc and two nitrogen of Phen-dione ligands. While, in Co-Fc, the octahedral geometry around Co([n] is satisfied by four sulfur atoms of two Fe-dtc ligands and two nitrogen atoms of Phen-dione ligand (Fig. 1). The Ni–S bond distances in Ni-Fc are 2.22 Å, while in Cu-Fc, the Cu–S distances are ~2.31 Å. The Ni–N, and Cu–N bond distances are 1.957 and 2.075 Å, respectively. In Co-Fc, the Co–S bond distance is ~2.31 Å and the Co–N bond distance is 2.020 Å. For Ni-Fc and Cu-Fc, the ∠S–Ni–S and ∠S–Cu–S are 79.06° and 77.48°, respectively. While ∠N–Ni–N and ∠N–Cu–N are 83.95° and 80.23°, respectively. In Co-Fc, the ∠S–Co–S and ∠N–Co–N bite angles are 51.48° and 81.44°, respectively. The geometrical parameters are in good agreement with the previously reported analogous complexes.37–45

The IR spectra of Ni-Fc, Cu-Fc, and Co-Fc displayed bands at 1583 cm⁻¹, 1512 cm⁻¹, and 1538 cm⁻¹, respectively, which can be attributed to the stretching frequencies of the C=O group of the 1,10-phenanthroline-5,6-dione ligand. The antisymmetric stretching frequencies corresponding to the –CS₂ moiety of dithiocarbamate ligand appears at 1195 cm⁻¹ (Ni-Fc), 1197 cm⁻¹ (Cu-Fc), and 1197 cm⁻¹ (Co-Fc). While, symmetric ν_CS₂ appears at 1037 cm⁻¹ (Ni-Fc), 1037 cm⁻¹ (Cu-Fc) and 1061 cm⁻¹ (Co-Fc). Also, ν_NCS₂ vibrations of the dithiocarbamate ligand appeared at 1430 cm⁻¹ (Ni-Fc), 1443 cm⁻¹ (Cu-Fc), and 1423 cm⁻¹ (Co-Fc) in all three sensitizers, which are intermediate between single (1360–1250 cm⁻¹) and double (1690–1640 cm⁻¹) stretching vibrations and hence indicates the existence of partial double bond character between carbon and nitrogen centers of the dithiocarbamate ligand.37–45

Electronic absorption spectroscopy

The electronic absorption spectroscopy on all three sensitizers was performed in DMF solutions (Fig. 2a). All three sensitizers display bands ~450 nm, which is due to the d → d (assigned to the 1Eₐg ← 1Aᵧg) transition of the ferrocene moiety.38 The higher energy absorptions observed near 360 nm, 300 nm and 250 nm in all the complexes may be assigned to the intraligand charge transfer transitions. Broad bands with medium intensity observed between 550–750 nm arise due to the square planar geometry around nickel([n] and copper([n] and the octahedral environment around cobalt([n]).38 Apart from this, the spectral response of the Co-Fc complex extends up to ~700 nm in comparison to Cu-Fc and Ni-Fc.

Also, the electronic absorption spectra of co-sensitizers Cu-Fc and Co-Fc with N719 dyes anchored on TiO₂ films were also recorded (Fig. 2b). The results indicated that after co-sensitization, the electronic absorption spectra of Cu-Fc and Co-Fc changed significantly. The electronic absorption responses of N719/Co-Fc, N719/Cu-Fc, and N719/Co-Fc/Cu-Fc in the visible region were enhanced, especially in the case of the triple sensitizer (N719/Cu-Fc/Co-Fc), it covered the spectral region from 400–720 nm. The enhancement in spectral response after co- and triple sensitization indicated that the light-harvesting ability of the DSSC devices fabricated using co- and triple sensitization strategies will be greatly improved, which may increase the short circuit current.

Electrochemical studies

One of the important prerequisites for the utilization of any sensitizer in a DSSC set-up is its energy level, which must be matching with respect to the conduction band of TiO₂ as it is important for the thermodynamic feasibility of the device. The sensitizers should have better regeneration efficiency and high electron injection efficiency for high-open-circuit voltage and...
short-circuit current of devices as well as high power conversion efficiency. Hence, in order to check the ability of electron injection and regeneration properties of sensitizers, cyclic voltammetry (CV) studies in a three-electrode cell on an electrochemical workstation, by using Pt disc as the working electrode and Ag/Ag⁺ (in acetonitrile) as the reference electrode, were performed. The electrochemical parameters of these complexes calculated from the CV are summarized in Table 1. From Table 1, it is evident that the LUMO level of sensitizers is more negative (−2.66 to −1.89 V vs. Ag/Ag⁺ in ACN) than the conduction band edge of the TiO₂ (−0.85 V vs. Ag/Ag⁺ in ACN). This indicates that the excited dyes can efficiently inject electrons into the conduction band of TiO₂ as well as the HOMO level of sensitizers is more positive (+0.146 to +0.152 V vs. Ag/Ag⁺ in ACN) than the redox potential of the electrolyte (−0.15 V Ag/Ag⁺ in ACN) providing an adequate driving force for regeneration of dyes (Fig. 3).

The energy level diagram (Fig. 4) obtained from the combined results of electronic absorption spectra and electrochemical data suggest a thermodynamically favourable electron injection from the excited state of N719 to the conduction band of TiO₂ followed by successive regeneration of N719 by an electrolyte (∙I⁻/I⁺). Also, the presence of co-sensitizers additionally transfers an electron to the conduction band of TiO₂ as well as the LUMO of N719 to uplift the photovoltaic performance with concomitant regeneration by the electrolyte.

Photovoltaic performance of DSSCs
The photovoltaic performances of the DSSC set-up fabricated using sensitization and co-sensitization strategies were assessed and the resultant photovoltaic parameters, obtained from the J–V curves of DSSCs (Fig. 5) are listed in Table 2. We can see that the highest photoelectric conversion efficiency (η) for the sensitization of the dyes in DSSC based on Co-Fc is 3.01%, with open circuit voltage (V_{oc}) 620 mV, short-circuit current (J_{sc}) 7.65 mA cm⁻², and fill factor (FF) 0.63, whereas the η for the DSSC based on Cu-Fc and Ni-Fc are 2.27% and 1.48%, respectively. Evidently, the high V_{oc} and J_{sc} for the Co-Fc-based DSSC are the key reasons for its high η. In comparison to Cu-Fc and Ni-Fc, Co-Fc exhibits better electron donation and conjugation properties due to the extended conjugation as well

| Sensitizer | E_{p,a} (V) | E_{p,c} (V) | ΔE = E_{p,a} − E_{p,c} (V) | E' = (E_{p,a} + E_{p,c})/2 (V) | E.S = E' + hν |
|------------|------------|------------|-----------------|-----------------|----------------|
| Co-Fc      | +0.178     | +0.126     | +0.052          | +0.152          | −1.89          |
| Cu-Fc      | +0.170     | +0.130     | +0.040          | +0.150          | −2.65          |
| Ni-Fc      | +0.167     | +0.125     | +0.042          | +0.146          | −2.66          |

Fig. 3 Cyclic voltammograms of the co-sensitizers recorded in DMF solution.

Fig. 4 Schematic energy level diagrams of HOMO and LUMO levels for the sensitizers and co-sensitizers compared to TiO₂.

Fig. 5 J–V curves for the DSSC cell set-up fabricated with different sensitizers and co-sensitizers.
as earlier absorption spectroscopic studies, also revealed that Co-Fc has a better light-harvesting ability in the visible region, which is favourable to increasing the $J_{oc}$ of the DSSCs. The spectral response of these dyes is not ideal to harvest the entire visible spectrum, so, for harvesting maximum visible light to the electrical energy, it is necessary to enhance the visible response of dyes. Hence, co-sensitization and double co-sensitization by combining N719 and one co-sensitizer (Cu-Fc or Co-Fc) and three dyes using N719 and two co-sensitizers (Cu-Fc and Co-Fc) to enhance the visible spectral response performed. Since the device performance of Ni-Fc was the poorest, this sensitizer was not used for co-sensitization studies further. By co-sensitizing, the spectral responses for Co-Fc/N719 and Cu-Fc/N719 were found to be extended in the visible region and it was from 450 nm to 750 nm and 400 nm to 650 nm for Co-Fc/N719 and Cu-Fc/N719, respectively, whereas, it covered the entire visible spectrum (400 to 750 nm) by double co-sensitization (Co-Fc/Cu-Fc/N719). As expected, the overall performance of the devices after co-sensitization and double co-sensitization improved significantly. Especially, for the DSSC based on Co-Fc/Cu-Fc/N719, the $J_{sc}$ was enhanced up to 12.87 mA cm$^{-2}$ with $V_{oc}$ and $\eta$ values were 733 mV and 6.05% respectively. From Table 2, it is evident that the enhancement in $\eta$ and $J_{sc}$ is consistent with the enhancement of their spectral performance. Further, we also found that the $V_{oc}$ of the DSSCs was improved considerably after co-sensitization and double co-sensitization, increasing by almost 100 mV. This enhancement in $V_{oc}$ is due to the compactness of dye molecule arrangement, which is favourable for inhibiting the penetration of the oxidizing electrolyte into the photoanode. Also, it helps to reduce the charge recombination behaviour. Also, an increase in short-circuit current density and the spectral sensitivity of devices provide a reinforcing effect on the device performances.

The incident photon-to-electron conversion efficiencies (IPCEs) for all the photovoltaic cell set-ups as a function of the wavelength of incident monochromatic light are presented in Fig. 6. The IPCE was determined from light-harvesting efficiencies, charge injection, charge regeneration, and charge collection properties. The sensitizers, Ni-Fc and Cu-Fc displayed maximum IPCE values of 42% and 44%, respectively at 440 nm, whereas Co-Fc exhibited maximum IPCE on higher wavelengths (500–750 nm) and it matches well with the electronic absorption spectrum of the individual dyes (Fig. 6, Table 2). In co-sensitized photovoltaic cell set-ups, superior IPCE values were obtained than those for sensitized cell set-ups. For co-sensitized cell set-ups, Co-Fc/N719, Cu-Fc/N719, and double co-sensitized cell set-ups Co-Fc/Cu-Fc/N719 the IPCE values ranged between 60 and 63% with the exposure of the entire visible region. The obtained higher IPCE value in co-sensitized than sensitized cell setups indicates a more efficient charge collection. Also, in comparison to the N719-based cell set-up, the photocurrent responses of Co-Fc/N719, Cu-Fc/N719, and Co-Fc/Cu-Fc/N719 were relatively better. The overall best performance of Co-Fc in sensitized and co-sensitized DSSC set-up was better due to the charge carrier separation and transportation with reduced electron–hole recombination. Also, the variation in the performance of sensitizers in sensitized and co-sensitized set-ups can also be correlated to their differences in the absorbance in the 400–500 nm and 600–750 nm regions, the spectral region where light absorption by N719 is inferior. The light absorption by sensitizers and co-sensitizers in this spectral region follows the order Co-Fc/Cu-Fc/N719 > Cu-Fc/N719 > Co-Fc/N719, which suggests that Co-Fc/Cu-Fc/N719 is most capable of recompensing the light absorption by N719 in 400–500 nm and 600–750 nm regions. In addition, the integration of the IPCE curves was accomplished by the convolution of the IPCE spectra where $J_{sc}$ for the DSSCs under solar illumination was predicted with the photo-flux density distribution for one sun as the following equation:

$$
J_{sc} = \int qE(\lambda)\text{IPCE}(\lambda)\,d\lambda
$$

where $q$ is the electron charge and $f(\lambda)$ is the incident photon flux density (air mass spectra [AM 1.5], ASTM G173) at wavelength $\lambda$. The $J_{oc}$ values from the IPCE curves of Co-Fc, Cu-Fc, Ni-Fc, Co-Fc/N719, Cu-Fc/N719, and Co-Fc/Cu-Fc/N719 are 6.86, 4.45, 3.18, 8.73, 10.32, 8.52 and 10.86 mA cm$^{-2}$, respectively, and match with the $J_{sc}$ values obtained from $J$–$V$ curves. The experiential differences between the two results are possible due to the spectral divergence and as the IPCE response, which was different from that of the simulated xenon lamp. Furthermore, desorption of dyes is common during co-sensitization, which may lead to a slight decline in photo-response current.

### Table 2

| Sensitizers | $J_{sc}$ (mA cm$^{-2}$) | $V_{oc}$ (mV) | FF | $\eta$ (%) | IPCE (%) |
|-------------|-------------------------|--------------|----|------------|----------|
| N719        | 9.68                    | 718          | 0.63 | 4.39       | 43       |
| Co-Fc       | 7.65                    | 620          | 0.63 | 3.01       | 44       |
| Cu-Fc       | 5.78                    | 616          | 0.64 | 2.27       | 44       |
| Ni-Fc       | 3.83                    | 610          | 0.63 | 1.48       | 42       |
| Co-Fc/N719  | 11.41                   | 730          | 0.64 | 5.31       | 62       |
| Cu-Fc/N719  | 10.67                   | 728          | 0.63 | 4.91       | 59       |
| Co-Fc/Cu-Fc/N719 | 12.87              | 733          | 0.64 | 6.05       | 63       |

**Electrochemical impedance spectroscopy (EIS)**

To assess the kinetics of the electron transport properties and charge recombination, electrochemical impedance
spectroscopy (EIS) was performed, which assists in the investigation of the charge transfer dynamics at the interfacial regions of the solid–liquid layers. The Nyquist plots of sensitized and co-sensitized photoelectrodes were recorded under standard global AM 1.5 solar irradiation in light and in dark at their open circuit potential (OCP) over a frequency range of $10^{-2}$ to $10^5$ Hz. In the EIS study, the semicircle in the middle frequency region is associated with the electron/charge transfer at the TiO$_2$/dye/electrolyte interface.

Fig. 7a and b show the Nyquist plots for DSSCs fabricated by using N719, Co-Fc/N719, Cu-Fc/N719 and Co-Fc/Cu-Fc/N719 in the dark and light, respectively. Under light illumination (Fig. 7b), the two semicircles located at high and middle-frequency regions are attributed to the electrochemical reaction at the Pt/electrolyte interface and charge transfer at the TiO$_2$/dye/electrolyte interface. The semicircles of large radii positioned at the middle frequency region in the Nyquist plots decrease after co-sensitization with the metal complex and the decline in radii of the semicircles are in the order Co-Fc/Cu-Fc/N719 < Co-Fc/N719 < Cu-Fc/N719 < N719. This signifies a decreasing trend of the electron transfer impedance ($R_{ct}$) as well as an increase in the charge transfer rate at this interface after co-sensitization. Under dark conditions (Fig. 7a), the appearance of the two semicircles in Nyquist plots at high and middle-frequency regions are attributed to the redox reaction at the Pt counter electrode and the electron transfer at the TiO$_2$/dye/electrolyte interface. In dark, the large semicircle lying in the middle frequency region represents the resistance of the charge transfer from TiO$_2$ to the electrolyte (back recombination resistance $R_{rec}$). The radii of the semicircles observed in the middle frequency range are in order of Co-Fc/Cu-Fc/N719 > Co-Fc/N719 > Cu-Fc/N719 > N719, indicating the sequence of $R_{rec}$ at the TiO$_2$/dye/electrolyte interface, which implies the retardation of the charge recombination between injected electrons and I$_3^-$ ions in the electrolyte, which as a consequence resulted in increased $V_{oc}$. The EIS study under light and dark conditions confirmed that the charge transfer rate increases and the charge recombination rate decreases for co-sensitized cell set-ups, which is conducive to enhancing the performance of DSSCs.

Table 3 summarises the DSSC results of the previously published co-sensitizers. According to the photovoltaic data, the performance of the co-sensitizers tested in this work is comparable to those of the previously published analogous co-sensitizers. However, the cell performance of the co-sensitizers is much lower than that of a few comparable derivatives, which may be attributed in part to the better spectrum responsiveness.
and photo-electron transfer. To achieve higher cell performance, co-sensitizers should display enhanced spectrum response in spectral regions where N719 exhibits poorer spectral response, better co-sensitizer anchoring, and effective photo-electron transfer to both the semiconductor conduction band and the excited state of N719.

Conclusion

In the presented investigation, three new ferrocene appended heteroleptic dithiocarbamate complexes with 1,10-phenanthroline-5,6-dione ancillary ligands were synthesized and used as co-sensitizers for state-of-the-art N719 dye. A new double co-sensitization strategy involving the use of Cu(n) and Co(III) complexes uplifted the photovoltaic performance of the state-of-the-art N719 dye and further uplifted DSSC performance by ~38% in comparison to the set-up constructed using only N719 dye under similar experimental condition. The enhancement in the photovoltaic performance of such a photovoltaic cell was accredited to the expansion of the electronic absorption band in the visible region and the rise in the electronic absorption intensity. Further, as evident by EIS studies, this improvement in cell performance was due to an increase in the charge transfer rate and a decline in the charge recombination rate after co-sensitization of N719. Such investigation will lay a firm platform to develop new co-sensitizers that along with N719 and analogous dyes will improve the DSSC performance.

Experimental

General considerations

All chemicals and reagents were obtained from commercial sources and used without any further purification. Elemental analyses for the co-sensitizers were performed on a PerkinElmer 240 C H N analyser, while FTIR spectra were recorded on a Shimadzu IR Affinity-1S spectrometer. The electronic absorption spectroscopy for the co-sensitizers was performed on a SPECORD 210 PLUS BU spectrophotometer. The mass spectrometry for all three co-sensitizers was performed on a Shimadzu IR A spectrometer. The electronic absorption spectroscopy for the co-sensitizers was performed on a PerkinElmer cation. Elemental analysis C14O2H8 (%): calcd, C, 80.76; H, 3.87. Found C, 81.12; H, 4.05. The obtained product was extracted using dichloromethane (3 x 50 mL) and washed with water followed by brine. The crude organic layer was dried over anhydrous sodium sulfate, filtered and again rotary evaporated to obtain an oily orange solid (N-ethanol-N-methylferrocenyl amine). The N-ethanol-N-methylferrocenyl amine ligand (0.251 g, 0.97 mmol) was added to it. After stirring for 30 min, Cs2 (0.114 g, 1.5 mmol) was added and the resultant reaction mixture was additionally stirred for 3 h to obtain a yellow precipitate, which was filtered and washed twice with diethyl ether and dried.

Characterization data. Yellow solid; yield: 0.243 g, 65.0%; mp 135–140 °C; 1H NMR (300 MHz, CDCl3, δ): 5.11 (s, 2H, Fe–CH3), 4.42 (s, 2H, Fe), 4.13 (s, 5H, Fe), 4.05 (s, 2H, Fe) 4.00 (t, 2H, J = 15.9 Hz, –CH2CH2OH), 3.55 (t, 2H, J = 18 Hz, –CH2CH2OH); 13C NMR (75.45 MHz, CDCl3, δ): 213.8 (CS2), 85.0 (Fe–C–C), 69.5 (C–H2), 68.0 (C–H3), 66.8 (C–H2), 58.9 (Fe–CH2), 52.5 (CH2–O), 51.0 (CH2–N); IR (KBr/cm−1, ν) 1438 (C–N), 1179 (C–S, anti-sym), 1063 (C–S, symm). Elemental analysis C14H16NFeS2K (%): calcd C, 47.37; H, 4.82; N, 4.05; S, 18.43.

Synthesis of 1,10-phenanthroline 5,6-dione: (Ph-dione)

Caution: Reaction led to the liberation of bromine fumes.

Potassium bromide (4.000 g, 33.6 mmol) and 1,10-phenanthroline (4.000 g, 22.2 mmol) were mixed in 200 mL two necked round bottom flask and to it was added the mixture of sulphuric acid (98%, 40 mL) and nitric acid (69%, 20 mL). During this addition, the temperature of the flask was maintained between 0–5 °C until the dark red-brown bromine fumes subsided and thereafter, the mixture was refluxed for 6 h. The obtained solution was cooled to room temperature and then poured over crushed ice. Thereafter, neutralisation of the reaction mixture using conc. NaOH solution between pH 4–5 yielded a yellow suspension, which was further extracted with chloroform (3 x 200 mL), washed with water (2 x 200 mL), and dried over anhydrous sodium sulfate, filtered and rotary evaporated to obtain a yellow residue. It was washed with diethyl ether and dried under a vacuum. The synthesis of 1,10-phenanthroline-5,6-dione,13a and [M] (Ph-dione)] M = Co, Ni, Cu was performed by considering the previously reported literature.43

Synthesis of potassium N-2-hydroxy-ethyl-N-methylferrocenyl dithiocarbamate

To the 10 mL ethanol solution of ferrocene carboxaldehyde (0.214 g, 1 mmol) was added ethanolamine (0.073 g, 1.2 mmol) and the reaction mixture was refluxed for 8 h under nitrogen to yield ferrocenyl imine derivative, which thereafter was reduced to a secondary amine with sodium borohydride (0.152 g, 4 mmol). The reaction mixture was rotary evaporated and the obtained product was extracted using dichloromethane (3 x 50 mL) and washed with water followed by brine. The crude organic layer was dried over anhydrous sodium sulfate, filtered and again rotary evaporated to obtain an oily orange solid (N-ethanol-N-methylferrocenyl amine). The N-ethanol-N-methylferrocenyl amine ligand (0.251 g, 0.97 mmol) was dissolved in THF/acetonitrile (50 : 50 v/v, 10 mL) and THF solution of KOH (0.067 g, 1.2 mmol) was added to it. After stirring for 30 min, Cs2 (0.114 g, 1.5 mmol) was added and the resultant reaction mixture was additionally stirred for 3 h to obtain a yellow precipitate, which was filtered and washed twice with diethyl ether and dried.

Characterization data. Yellow solid; yield: 0.243 g, 65.0%; mp 135–140 °C; 1H NMR (300 MHz, CDCl3, δ): 5.11 (s, 2H, Fe–CH3), 4.42 (s, 2H, Fe), 4.13 (s, 5H, Fe), 4.05 (s, 2H, Fe) 4.00 (t, 2H, J = 15.9 Hz, –CH2CH2OH), 3.55 (t, 2H, J = 18 Hz, –CH2CH2OH); 13C NMR (75.45 MHz, CDCl3, δ): 213.8 (CS2), 85.0 (Fe–C–C), 69.5 (C–H2), 68.0 (C–H3), 66.8 (C–H2), 58.9 (Fe–CH2), 52.5 (CH2–O), 51.0 (CH2–N); IR (KBr/cm−1, ν) 1438 (C–N), 1179 (C–S, anti-sym), 1063 (C–S, symm). Elemental analysis C14H16NFeS2K (%): calcd C, 47.37; H, 4.82; N, 4.05; S, 18.43.
**Characterization data [Co(Phen-dione)].** Olive brown solid; yield: 0.175 g, 51.62%; mp 182–187 °C (dec). Elemental analysis C_{12}H_{5}Cl_{2}CoN_{2}O_{3} (%): calcd, C, 42.51; H, 1.47; N, 8.15. Found C, 42.89; H, 1.73; N, 8.34.

**Characterization data [Ni(Phen-dione)].** Brown red solid; yield: 0.175 g, 55%; mp 178–181 °C (dec). Elemental analysis C_{12}H_{5}Cl_{2}NiN_{2}O_{3} (%): calcd, C, 41.94; H, 1.47; N, 8.34. Found C, 41.98; H, 3.08; N: 5.87; S, 9.03. IR (KBr/cm\(^{-1}\), ν) 3066 (Ar–C–H), 1538 (C–O), 1197 (C–S(antisymmetric)), 1037 (C–S(symm)), 622 (Fe–C, Cp ring). HRMS (ESI): m/z = 605.9707 [M – 2H]\(^+\); for [FeC_{26}H_{20}O_{4}N_{4}S_{2}Co]^{+} 605.9922.

**Synthesis of [Co(Fcdtc)\(_2\)(Phen-dione)]PF\(_6\) (Co-Fc).** A methanolic solution of KPF\(_6\) (0.184 g, 1 mmol) was added dropwise to the stirring solution of Fcdtc (0.186 g, 0.5 mmol). The obtained light yellow solution was stirred for half an hour in an ice bath followed by the addition of [Co(Phen-dione)] (0.169 g, 0.5 mmol). Upon complete addition, the colour of the reaction mixture turned olive brown and the reaction mixture was stirred overnight. The solvent was rotary evaporated and the obtained residue was washed with petroleum ether and dried under a vacuum.

**Characterization data.** Olive brown solid; yield: 0.175 g, 69.5%; mp 194–200 °C (dec). Elemental analysis Fe_{2}C_{40}H_{38}O_{2}S_{4}N_{3}PF_{6} (%): calcd, C, 41.47; H, 2.95; N, 5.58; S, 8.52; found C, 41.98; H, 3.08; N, 5.87; S, 9.03. IR (KBr/cm\(^{-1}\), ν) 3066 (Ar–C–H), 1538 (C–O), 1197 (C–S(antisymmetric)), 1037 (C–S(symm)), 622 (Fe–C, Cp ring). HRMS (ESI): m/z = 605.9707 [M – 2H]\(^+\); for [FeC_{26}H_{20}O_{4}N_{4}S_{2}Co]^{+} 605.9922.

**Fabrication of DSSCs**

For the fabrication of DSSC, FTO (F-doped SnO\(_2\) purchased from Pilkington. Co. Ltd., 8Ω/√\(\text{cm}\)) glass plates were cleaned in ethanol by using an ultrasonic bath for 30 min. FTO was used to prepare both the photo- and counter-electrodes. The TiO\(_2\) thin films were prepared using a commercial TiO\(_2\) paste (Ti-Nanoxide T, Solaronix) and using the doctor-blade technique onto the FTO substrate followed by calcination at 450 °C for 30 min. The sensitized and co-sensitized photoelectrodes were prepared by immersing the thin nanoporous TiO\(_2\) film (~6 μm) in a 1 : 1 ratio of DMF: ethanol solution of Co-Fc, Cu-Fc, Ni-Fc, and 1 : 1 ratio of DMF: ethanol solution of Cu-Fc/N719, Cu-Fc/N719, and Co-Fc/Cu-Fc/N719 correspondingly for 6 h and then washing with dichloromethane, followed by ethanol and drying at room temperature. The platinum counter-electrodes were prepared by screen printing of the Pt catalyst (Solaronix) on FTO and then sintering at 400 °C for 30 min. The dye-anchored TiO\(_2\) photoelectrodes and Pt counter-electrodes were assembled into face to face sandwich type assembly, which was sealed from three sides by using a hot-melt sealant film (Solaronix) as a spacer between the electrodes by heating it at 80 °C. A drop of electrolytic solution (0.05 M iodine, 0.05 M I\(_2\), and 0.5 M 4-tert butylpyridine in acetonitrile) was placed on the open side of the cell assembly, which was driven inside the cell by capillary action and then the open side was closed by using Araldite followed by copper connection on both the electrodes with the help of silver paste and Araldite.

**Characterization of DSSCs**

The photoelectrochemical performance characteristics (short circuit current \(I_{sc}\) (mA cm\(^{-2}\)), open-circuit voltage \(V_{oc}\) (V), fill factor FF, and overall conversion efficiency \(η\)) were measured under illumination with a 1000 W xenon lamp (Oriel 91193) using a Keithley Model 2400. The light intensity was confirmed to be homogenous over an \(8 \times 8\) in\(^2\) area by calibration with a Si solar cell for 1 sunlight intensity (AM 1.5G, 100 mW cm\(^{-2}\)). An accidental increase in the temperature inside the cell was prevented using a cooler with a propeller. Incident photon-to-current conversion efficiency (IPCE) for the dyes was measured as a function of wavelength from 400–700 nm (PV Measurement Inc.) using a standard tungsten–halogen lamp as monochromatic light and a broadband bias light for approximating 1 sunlight intensity. The electrochemical impedance spectroscopy (EIS) was performed on a CH 660C electrochemical analyzer (CH Instruments, Shanghai, China) in a two-
electrode configuration. The photoanode was used as a working electrode and the Pt electrode as a counter electrode. The electron transport properties were investigated using electrochemical impedance spectroscopy (EIS) with a 10 mV alternation signal in the frequency range of $10^{-2}$ to $10^5$ Hz. UV-vis absorption spectra were recorded on a Shimadzu UV-3600 spectrometer. Cyclic voltammetry (CV) was performed using absorption spectra were recorded on a Shimadzu UV-3600 spectrometer. The amount of dye loading on the TiO$_2$ electrode was measured using a Shimadzu UV-3600 spectrometer. The sensitized electrodes were immersed in a 0.1 M NaOH solution in a mixed solvent 1 : 1 of DCM : ethanol, which resulted in the desorption of each dye. The dye loading amount can be estimated by using formula: $C = AV/eS_o$ (where $C$ is the amount of dye loading, $A$: is the optical absorbance of the dye, $V$: is the volume of desorption solution, $e$: is the molar extinction coefficients and $S_o$: effective area of TiO$_2$ films).

Computational details
In order to assess the structures of all three sensitizers, density functional theory calculations were executed. The molecular geometries of all the three compounds were optimized using the B3LYP exchange-correlation functional. The MDF10 basis set for Fe, Co, Ni, and Cu was used, while the 6-31G** basis set for other atoms was used for geometry optimization. All calculations were performed using the Gaussian 09 program.

Conflicts of interest
The authors declare no conflicts of interest.

Acknowledgements
Dr Mohd. Muddassir is grateful to Researchers Supporting Project number (RSP-2021/141), King Saud University, Riyadh, Saudi Arabia, for financial assistance. AK is grateful to the Research and Development, Department of Higher Education, UP Government for sanctioning project number 108/2021/2585/sattar-4-2021-4(28)/2021.

References
1 D. Raja, B. Selvaraj, G. Shanmugam, A. Maruthapillai and D. Sundaramurthy, *Energy Fuels*, 2021, 35, 4273–4282.
2 A. Y. Hoekstra and T. O. Wiedmann, *Science*, 2014, 344, 1114–1117.
3 X.-B. Li, C.-H. Tung and L.-Z. Wu, *Nat. Rev. Chem.*, 2018, 2, 160–173.
4 C. Dragonetti and A. Colombo, *Molecules*, 2021, 26, 2461.
5 M. Kokkonen, P. Talebi, J. Zhou, S. Asgari, S. A. Soomro, F. Elsehrawy, J. Halme, S. Ahmad, A. Hagfeldt and S. G. Hashmi, *J. Mater. Chem. A*, 2021, 9, 10527–10545.
6 B. O’Regan and M. Grätzel, *Nature*, 1991, 353, 737–740.
7 (a) M. Grätzel, *Nature*, 2001, 414, 338–344; (b) H. L. Jia, S. S. Li, B. Q. Gong, L. Gu, Z. L. Bao and M. Y. Guan, *Sustainable Energy Fuels*, 2020, 4, 347–353.
8 V. K. Singh, R. K. Kanaparthi and L. Giribabu, *RSC Adv.*, 2014, 4, 6970–6984.
9 P. S. Gangadhar, A. Jagadeesh, A. S. George, G. Reddy, S. Prasanthkumar, S. Soman and L. Giribabu, *Mol. Syst. Des. Eng.*, 2021, 6, 779–789.
10 M. V. Vinayak, T. M. Lakshmykanth, M. Yoosuf, S. Soman and K. R. Gopidas, *Sol. Energy*, 2016, 124, 227–241.
11 K. Kakiage, Y. Aoyama, T. Yano, K. Oya, J.-I. Fujisawa and M. Hanaya, *Chem. Commun.*, 2015, 51, 15894–15897.
12 L. Giribabu, R. K. Kanaparthi and V. Velkannan, *Chem. Rec.*, 2012, 12, 306–328.
13 Q. Yu, Y. Wang, Z. Yi, N. Zu, J. Zhang, M. Zhang and P. Wang, *ACS Nano*, 2010, 4, 6032–6038.
14 Z.-S. Wang, Y. Cui, Y. Dan-oh, C. Kasada, A. Shinpo and K. Hara, *J. Phys. Chem. C*, 2007, 111, 7224–7230.
15 S. Ferrere and B. A. Greg, *New J. Chem.*, 2002, 26, 1155.
16 T. Horiuchi, H. Miura and S. Uchida, *Chem. Commun.*, 2003, 24, 3036–3037.
17 K. Devulapally, G. Reddy, S. Prasanthkumar, A. Jagadeesh, S. Soman and L. Giribabu, *J. Porphyryns Phthalocyanines*, 2021, 25, 407–417.
18 J. V. S. Krishna, D. Koteswar, T. H. Chowdhury, S. P. Singh, I. Bedja, A. Islam and L. Giribabu, *J. Mater. Chem. C*, 2019, 7, 13594–13605.
19 V. Nikolov, A. Charisiadis, S. Chalkiadaki, I. Alexandropoulos, S. C. Pradhan and S. Soman, *Polyhedron*, 2018, 140, 9–18.
20 V. K. Narra, H. Ullah, V. K. Singh, L. Giribabu, S. Senthilarasu, S. Z. Karazhanov, A. A. Tahir, T. K. Mallick and H. M. Upadhyaya, *Polyhedron*, 2015, 100, 313–320.
21 M. Urbani, M. Grätzel, M. K. Nazeeruddin and T. Torres, *Chem. Rev.*, 2014, 114, 12330–12396.
22 S. Cherian and C. C. Wamser, *J. Phys. Chem. B*, 2000, 104, 3624–3629.
23 W. M. Campbell, K. W. Jolley, P. Wagner, K. Wagner, P. J. Walsh, K. C. Gordon, L. Schmidt-Mende, M. K. Nazeeruddin, Q. Wang and M. Grätzel, *J. Phys. Chem. C*, 2007, 111, 11760–11762.
24 A. Yella, H.-W. Lee, N. Tsao, C. H. Yi, A. K. Chandiran, M. K. Nazeeruddin, E. W.-G. Diao, C.-Y. Yeh, S. M. Zakeeruddin and M. Grätzel, *Science*, 2011, 334, 629–634.
25 S. Mathew, A. Yella, P. Gao, R. Humphry-Baker, B. F. E. Curchod, N. Ashari-Astani, I. Tavernelli, U. Rothlisberger, M. K. Nazeeruddin and M. Grätzel, *Nat. Chem.*, 2014, 6, 242–247.
26 J.-M. Ji, H. Zhou, Y. K. Eom, C. H. Kim and H. K. Kim, *Adv. Energy Mater.*, 2020, 10, 2000124.
27 D. Koteswar, S. Prashathkumar, S. P. Singh, T. H. Chaudhar, I. Bedja, A. Islam and L. Giribabu, *Mater. Chem. Front.*, 2022, 6, 580–592.
28 Y. K. Eom, S. H. Kang, I. T. Choi, Y. Yoo, J. Kim and H. K. Kim, *J. Mater. Chem. A*, 2017, 5, 2297–2308.
29 N. V. Krishna, J. V. S. Krishna, S. P. Singh, L. Giribabu, L. Han, I. Bedja, R. K. Gupta and A. Islam, *J. Phys. Chem. C*, 2017, 121, 6464–6477.
