Review

Azaacenes Based Electroactive Materials: Preparation, Structure, Electrochemistry, Spectroscopy and Applications—A Critical Review†

Kamil Kotwica1,*, Ireneusz Wielgus2 and Adam Pron2

Abstract: This short critical review is devoted to the synthesis and functionalization of various types of azaacenes, organic semiconducting compounds which can be considered as promising materials for the fabrication of n-channel or ambipolar field effect transistors (FETs), components of active layers in light emitting diodes (LEDs), components of organic memory devices and others. Emphasis is put on the diversity of azaacenes preparation methods and the possibility of tuning their redox and spectroscopic properties by changing the C/N ratio, modifying the nitrogen atoms distribution mode, functionalization with electroaccepting or electrodonating groups and changing their molecular shape. Processability, structural features and degradation pathways of these compounds are also discussed. A unique feature of this review concerns the listed redox potentials of all discussed compounds which were normalized vs. Fc/Fc+ This required, in frequent cases, recalculation of the originally reported data in which these potentials were determined against different types of reference electrodes. The same applied to all reported electron affinities (EAs). EA values calculated using different methods were recalculated by applying the method of Sworakowski and co-workers (Org. Electron. 2016, 33, 300–310) to yield, for the first time, a set of normalized data, which could be directly compared.

Keywords: azaacenes; synthesis; functionalization; redox properties; devices

1. Acenes vs. Azaacenes—Generalities

In the past two decades significant research effort in the domain of organic semiconductors was directed towards acenes and azaacenes [1–4]. Several types of acenes were tested as semiconducting active layers in p-channel organic field effect transistors (OFETs). Interesting properties of these compounds resulted from a relatively low value of their ionization potential which facilitated the injection of holes to the active layer in the transistor operating conditions. Investigated acenes readily crystallized due to planarity and frequent high symmetry of their molecules, yielding good quality single crystals or highly crystalline thin layers when deposited on an appropriate substrate. Some of them exhibited extremely high value of the hole mobility. For example, hole mobility of 40 cm²/Vs was reported for single crystals of rubrene (5,6,11,12-tetraphenyltetracene) [5].

From the point of view of technological applications, limited stability of acenes towards oxidative degradation constitutes the main drawback of these semiconductors. This especially applies to higher acenes consisting of several aromatic rings. Formation of carbonyl groups as a result of oxidative degradation perturbs the conjugation in individual molecules and induces disorder in their supramolecular organization, resulting in a drastic decrease of the charge carriers mobility [6].
Acenes cannot be used as semiconductors transporting electrons in any organic electronic devices, including n-channel OFETs, because their LUMO level is too high, implying low electron affinity (EA) values and by consequence very difficult electron injection [7,8].

Azaacenes (N-heteroacenes) are derivatives of acenes in which a fraction of carbon atoms are substituted by more electronegative nitrogen atoms. This substitution induces significant changes in their electronic, redox and spectroscopic properties. Quantum-chemical calculations, and especially those based on Density Functional Theory (DFT), constitute a powerful tool for the elucidation of the effect of N/C ratio and the N atoms distribution mode on the electronic and redox properties of azaacenes. This approach allows for the determination of their HOMO and LUMO levels as well as their ionization potentials (IPs) and electron affinities (EAs) in vacuum or in a given solvent. The latter can be confronted with experiments consisting of chronovoltamperometric determination of IP and EA. Even though the experimental values may sometimes differ from the calculated ones, especially in the case of EA, the observed trends usually remain the same.

As demonstrated by DFT calculations, for linear acenes and azaacenes the positions of HOMO and LUMO levels are strongly dependent on the number of aromatic rings in the molecule. In particular, LUMO level decreases and HOMO level increases with increasing length of the molecule [9]. If acene and azaacene of the same number of rings are compared, both HOMO and LUMO levels are lowered with increasing N/C ratio. For example, LUMO and HOMO levels of pentacene are located at $-2.7$ eV and at $-4.6$ eV while in the corresponding diazapentacene (naphtho[2,3-b]phenazine) these levels are lowered to $-3.3$ eV and $-5.1$ eV, respectively. In tetrazapentacene (quinoxalino[2,3-b]phenazine) they are further lowered to $-3.6$ eV and $-5.3$ eV [10]. Moreover, in the case of azaacenes of the same number of rings and the same number of nitrogen atoms, the HOMO and LUMO energies are sensitive to the nitrogen atoms distribution. Nitrogens located in terminal rings have less pronounced effect on the energies of LUMO and HOMO than those from inner rings. Similar trends are observed in the case of azahexacenes [11,12]. HOMO and LUMO energies as well as closely related to them through the Koopmans theorem IP and EA values determine electronic, spectroscopic and redox (electrochemical) properties of azaacenes.

Trends evidenced by DFT calculations are corroborated by experiment. In Table 1 experimentally determined redox potentials, electron affinities and optical band gaps are compared for the shortest acenes (naphthalene and anthracene) and their azaacene analogues containing increasing number of nitrogen atoms.

Table 1. Reduction potentials, electron affinities and optical band gaps of selected acenes and their azaacene analogues.

| Compound          | $E_{1/2 \text{red}}$ vs. Fe/Fe$^+$ [V] | EA * [eV] | $E_{\text{g opt}}$ [eV] |
|-------------------|---------------------------------------|-----------|-------------------------|
| **Acenes**        |                                       |           |                         |
| Naphthalene       | $-3.44$ [13]                          | 0.77      | 4.14 [14]               |
| Anthracene        | $-2.91$ [15]                          | 1.40      | 3.20 [14]               |
| **Azaacenes**     |                                       |           |                         |
| Quinoline (1N)    | $-2.77$ [15]                          | 1.56      | 3.94 [16] **            |
| Quinoxaline (2N)  | $-2.26$ [15]                          | 2.16      | 3.85 [17] **            |
| Acridine (1N)     | $-2.26$ [15]                          | 2.16      | 3.22 [18] **            |
| Phenazine (2N)    | $-1.83$ [15]                          | 2.67      | 3.07 [19] **            |
| Pyrazinoquinoxaline (4N) | $-1.60$ [20] | 2.94 | 2.95 [21] ** |

* recalculated from original data in which potential was given vs. SCE. * EA calculated by following the procedure proposed in [22]. ** $E_{\text{opt}}$ estimated from the absorption spectra shown in reference [16–19,21] by determining the wavelength of the absorption onset and converting it from nm to eV.
As evidenced from these data, the band gap ($E_{g\text{opt}}$) decreases with increasing N/C ratio whereas the electron affinity (EA) increases in full accordance with DFT data. In particular, the reduction potential of naphthalene is significantly lower than the corresponding potentials of its aza analogues, i.e., quinoline and quinoxaline. The same applies to anthraquinone and its aza-derivatives. Note that for the same number of aromatic rings (three) EA increases in a series: acridine (1N), phenazine (2N), pyrazinoquinoxaline (4N). This trend is maintained in azaacenes of higher molecular mass and different molecular shape.

Further tuning of electronic, optical and redox properties of azaacenes is possible by introduction of substituents of electrodonating or electroaccepting properties. Solution processability of azaacenes, frequently very difficult or even nonexistent, can also be improved by attaching appropriate solubilizing groups to the conjugated core.

Azaacenes are aromatic compounds, however their expected aromaticity should be lower than that of the corresponding acenes. It is instructive to compare aromaticities of the simplest aromatic and aza-aromatic compounds such as benzene, pyridine and three diazines, i.e., 1,2-diazine (pyridazine), 1,3-diazine(pyrimidine) and 1,4-diazine (pyrazine). Pyridine, pyrazine and pyrimidine are highly aromatic. Their HOMA (Harmonic Oscillator Model of Aromaticity) indices exceed 0.99. The only exception is pyridazine (1,2-diazine) whose HOMA index is significantly lower and equals to 0.911 [23]. This pronounced decrease of aromaticity is caused by the presence of two nitrogen atoms at adjacent positions in the ring, forming an N-N bond. Acenes and azaacenes reflect this trend indicating higher aromaticity of the former. It should also be noted that their HOMA indices are in line with those derived from NICS (Nucleus-Independent Chemical Shifts) [24].

Redox properties of azaacenes are inherently associated with their chemical constitution involving aromatic core and eventually the presence of electron donating (electron accepting) substituents. Redox properties of acenes and azaacenes are most frequently experimentally determined by cyclic voltammetry (cv) or differential pulse voltammetry (dpv). These data are usually available in literature; however, the determined redox potentials are expressed versus different reference electrodes which makes their comparison difficult. For comparative reasons all redox potentials presented in this review are expressed vs. ferrocene/ferrocenium couple ($Fc/Fc^+$). This means that all redox potentials reported versus different than $Fc/Fc^+$ references were recalculated by the authors of this review. This approach allows for direct comparison of the redox properties of all azaacenes discussed here.

The electrochemically measured redox potentials can in principle be used for the determination of the solid state ionization energies such as ionization potential (IP) or electron affinity (EA) of electroactive molecules, i.e., parameters which are crucial for the majority of organic semiconductors applications. This approach requires, however, some comments. IP and EA can be directly measured on thin organic semiconductor films using ultraviolet photoelectron spectroscopy (UPS) and inverse photoelectron spectroscopy (IPES), respectively. Electrochemical experiments are carried out in solution, thus in this respect they do not provide direct access to IP and EA. In the majority of papers IP and EA are calculated from the electrochemical data using the following equations:

\[
\text{EA} = e\text{E}_{\text{red}} + C
\]

In addition:

\[
\text{IP} = e\text{E}_{\text{ox}} + C
\]

where $E_{\text{red}}$ and $E_{\text{ox}}$ stand for the first reduction potential and the first oxidation potential of the studied molecule, respectively; C is the absolute energy of the $Fc/Fc^+$ redox couple in the electrolyte solution.

The value of C is still the subject of debate and its value can range from 4.8 eV to 5.1 eV in various reports (see for example discussion in [25]). Applications of different values of C makes the EA and IP values, reported in various papers, difficult to compare.
Moreover, as pointed out by Sworakowski et al. in a series of papers [22,26,27], the EA and IP values derived from electrochemical experiments carried out in solutions and calculated using Equations (1) and (2) may significantly differ from those determined for thin films by UPS and IPES, thus not giving direct access to solid state ionization energies. These authors, analyzing all available electrochemical and photoelectron spectroscopy data, pointed out that a clear correlation exist between the electrochemically determined redox potentials and experimentally determined IP and EA values which can be expressed by the following equations:

\[ EA = 1.18 \times eE_{\text{red}} + 4.79 \text{ [eV]} \] (3)

In addition:

\[ IP = 1.15 \times eE_{\text{ox}} + 4.83 \text{ [eV]} \] (4)

Again, for comparative reasons we used Equations (3) and (4) for the calculations of EA and IP values for all azaacenes reported in this review.

Depending on N/C ratio azaacenes can show either n-type, p-type or ambipolar character in the field effect transistor configuration. The latter means that they can transport either holes or electrons, depending on the electrodes’ polarity. For high N/C ratio they lose their ambipolar character and are used as electrons transporting materials in n-channel field effect transistors. This is expected, since a significant increase of EA of these compounds facilitates the stabilization of radical anions formed temporarily upon the injection of electrons from the source electrode [28,29]. An increase of IP of azaacenes as compared to acenes leads to a significant improvement in their resistance against oxidative degradation, making the formation of carbonyl groups and/or degradative dimerization more difficult [30]. Azaacenes find their application as organic semiconductors not only in electronics but also in optoelectronics as electroluminesphores in organic light emitting diodes (OLEDs) [31–34] and in photovoltaics as acceptors in non-fullerene organic photovoltaic cells [35–37].

Research on electroactive azaacenes is principally focused on three types of compounds: (i) linear which can be considered as analogues of the corresponding linear acenes; (ii) more condensed azaacenes containing pyrene-type segments; (iii) star-shaped azaacenes. This classification should be considered as somehow fuzzy since large number of azaacenes exist which cannot be attributed to any of these groups or they are on the borderline of two groups.

2. Synthesis of Azaacenes

First syntheses of azaacenes were reported at the end of 19th century by Fischer and Hepp [38,39]. These research efforts were then continued through the 20th century and significantly intensified in recent years [40,41].

Initially elaborated methods of azaacenes synthesis were based on reactions of o-dihalogeno (or o-dihydroxyaryl) aromatic compounds with aromatic o-diamines. The condensation reactions were usually carried out at temperatures exceeding melting points of the substrates, typically in the range of 130–160 °C, but sometimes at temperatures exceeding 200 °C. In these reaction conditions the target azaacenes (or dihydroazaacenes) could be obtained in reasonable yields. The above outlined simple preparation methods are exploited even nowadays, especially in cases where the substrates and the product are insufficiently soluble in conventional organic solvents [42]. Low reactivity of aryl o-diamines can be considered as a somehow weaker point of this synthetic pathway [43].

Over the years, the condensation methods of azaacenes preparation have been constantly modified and improved. In particular, application of protic compounds as solvents or catalytic admixtures led to a significant lowering of the reaction temperature to values below 100 °C (in selected cases even below 50 °C), while retaining high reaction yields [44–46].

Condensation of aromatic o-diamines with o-dihydroxy derivatives of aryl compounds carried out in mild conditions yields dihydroazaacenes whereas the condensation with
o-diketones results in azaacenes. In the case of substrates containing hydroxyl groups in addition to keto ones, as in the synthesis with the use of dihydroxybenzoquinone or oxalic acid, dihydroxy derivatives of azaacenes are obtained in the first step, then dihydroazaacenes in the second one, which finally have to be oxidized to tetraazaacenes [45,46].

Optimization of o-diketone condensation with o-diamines (or their salts) leads to various pyrazine derivatives which can be obtained in good yields. Various azaacenes of extended structures were obtained at room temperatures using acetic acid in chloroform [47]. It should be noted that azaacenes of very diversified nitrogen atoms distribution can be obtained via condensation of amines with ketones. For example, Zhu et al. [48] prepared phthalazine derivatives in which nitrogen atoms occupied adjacent positions.

An interesting procedure of the synthesis of 1,4,5,8-tetraazaanthracene functionalized with four solubility-inducing ester groups was reported [49]. In this method, protected substrate, namely 1,2,4,5–benzene tetraamine tetrahydrochloride, upon deprotection in the reaction medium with a weak base (sodium acetate), reacted with dioxosuccinic acid ester to yield the desired product (see Figure 1). The reaction yield was 50%. Further modification through trans-esterification allowed for introduction of R groups different from the initial ones.

![Figure 1. Preparation of 1,4,5,8-tetraazaanthracene containing four solubilizing ester groups, R = Me, Et or Bu.](image)

The most general approach to the synthesis of hydroazaacenes, which upon oxidation can be transformed into azaacenes, is based on the Buchwald-Hartwig coupling reaction. This especially applies to dihydrophenazines but also to longer hydroazaacenes in a form of tetrahydroazaacenes or hexahydroazaacenes [50–52]. Buchwald-Hartwig coupling allows for the formation of C-N bonds in significantly milder conditions as compared to other methods, facilitating in this manner the introduction to the conjugated core substituents of different type. In the simplest version of the discussed reaction aryl o-diamines are reacted with aryl o-dihalides in the presence of a palladium complex catalyst and a base. Ligands in the palladium complex are appropriately selected, depending on the solvent used (see Figure 2).

![Figure 2. General scheme of Buchwald-Hartwig coupling, Br or Cl.](image)

An instructive approach to the preparation of azaacenes via Buchwald-Hartwig coupling can be found in reference [43].

Reaction yields in Buchwald-Hartwig coupling depend on the nature of substituents in the substrates [50,51]. In particular, in the synthesis of dihydrophenazine derivatives, a decrease of the reaction yield was observed if substituents showing electron accepting properties were introduced to 2,3-dichloroquinoxaline. Functionalization of diamine substrates had less pronounced effect on the obtained reaction yields. Similar tendency was also observed in the synthesis of fluoro derivatives of azapentacenes which were obtained with a significantly lower yields as compared to their non-fluorinated counterparts [53].

One has to be aware of the fact, that in this reaction two C-N bonds have to be formed in the ortho positions with respect to each other. This may induce steric problems which
then, together with lower reactivity of o-dihalides containing electron accepting groups, may in some cases result in lowering the reaction yield [54].

In view of these briefly discussed shortcomings of Buchwald-Hartwig coupling in the synthesis of azaacenes containing strongly electron accepting substituents, alternative approaches were proposed. Schwaben et al. [53] obtained fluoro-substituted diazapentacenes from aromatic o-diamines and perfluorinated benzene or naphthalene using nucleophilic substitution (SNAr). In the first step the treatment with NaH or with sodium bis(trimethylsilyl) amide at RT resulted in the formation of a C-N bond. In the second step, carried out at elevated temperature for extended time, dihydropyrazine ring closure occurred in the presence of a weak base (i-Pr2EtN) to yield the target tetra-fluorodiazatetracene. Comparative studies of Buchwald-Hartwig coupling and nucleophilic substitution showed that in some cases the latter resulted in better yields and enabled syntheses of azaacene derivatives unachievable by the former.

In the preparation using Buchwald-Hartwig coupling or related processes, hydroazaacenes are frequently obtained as intermediate products, which have to be then oxidized to yield the target azaacenes. Selection of an appropriate oxidant is of crucial importance in this case. Manganese dioxide (MnO2) in chloroform or dichloromethane is one of the most popular oxidizing agents used in the preparation of various types of diazaacenes or tetraazaacenes containing deactivating halo-, nitro- and other substituents [50,52,55]. It was also used in the synthesis of hexaazaacenes [51]. Other oxidants were also applied with success such as N-bromoimide of succinic acid, 2-iodobenzoic acid, copper (II) acetate, potassium dichromate, pyridine dichromate, pyridine chlorochromate, 2,3-dichloro-5,6-dicyano-p-benzoquinone, chloranil and lead (IV) dioxide. PbO2 readily converts hydroazaacenes into azaacenes but in the same time it does not oxidize alkylsilyl groups which are frequently attached to the azaacene core with the goal to improve their solution processability and making them more compatible upon deposition on silicon substrates in hybrid organic/inorganic electronic devices [56]. Oxidation of hydroazaacenes to azaacenes has to be performed with caution because in the case of azaacenes of low N/C ratios, too strong oxidants may result in the formation of benzoquinone-type units, significantly worsening semiconducting properties of these compounds.

An interesting procedure of the preparation of azaacenes was proposed by Raju et al. [57]. These authors developed one-pot synthesis of benzophenazine and its derivatives by reacting benzoxepine-4-carboxylates with benzene-1,2-diamines, in the presence of Bi(OTf)3 catalyst. The reaction, carried out under relatively mild conditions resulted in very good yields.

Exploiting appropriate catalytic systems, it is possible to prepare azaacene derivatives in which the distribution of nitrogen atoms differs from that characteristic of the 1,4-pyrazine unit. Reference [58] can be considered here as an instructive example. These authors described the application of nickel-based catalysts in the synthesis of quinazoline derivatives containing either aryl or alkyl substituents in position 2. In this procedure benzylamine with unoccupied ortho position was reacted with nitryl group containing benzene, pyridine or another appropriate aromatic reagent. The resulting azaacene was substituted in position 2 with either aliphatic or aromatic substituent originating from the nitryl derivative. This promising method should lead to quinazoline derivatives of more extended aromatic structure.

Kotwica et al. [59,60] developed a very simple method of the preparation of phenazine derivatives. These authors demonstrated that indanthrone (6,15-dihydridinaphtho[2,3-a:2′,3′-h]phenazine-5,9,14,18-tetraone) an old, nitrogen containing insoluble dye can be converted into a solution processable azaacene, namely tetraoctyloxydinaphtho[2,3-a: 2′,3′-h]phenazine, in a simple one-pot process consisting of carbonyl groups reduction with sodium dithionite, followed by o-alkylation under phase transfer catalysis conditions (see Figure 3a). Continuing their research on nitrogen-containing vat dyes as substrates for the preparation of azaacenes Kotwica et al. [61,62] prepared a series of solution processable fused azaacenes, namely 8,16-dialkoxybenzo[h]benz[5,6]acridino[2,1,9,8-klmna]acridines.
from flavanthrone (benzo[h]benz[5,6]acridino[2,1,9,8-klmna]acridine-8,16-dione)—an almost forgotten dye (see Figure 3b).

![Figure 3](image1)

**Figure 3.** Preparation of: (a) tetraalkoxydipnaphtho[2,3-a: 2′,3′-h]phenazine from 6,15-dihydodinaphtho-[2,3-a:2′,3′-h]phenazine-5,9,14,18-tetraone (flavanthrone) R = n-alkyl; (b) 8,16-dialk oxybenzo-[h]benz[5,6]acridino[2,1,9,8-klmna]acridines from benzo[h]benz[5,6]acridino[2,1,9,8-klmna]acridine-8,16-dione. R = n-alkyl group.

Very similar phenazine derivatives were obtained by oxidation of dialkyl-substituted 2-aminoanthraquinone with 2,3-dichloro-5,6-dicyanobenzoquinone [63]. Depending on the solvent used, two products in varying proportions were obtained: Z-shaped phenazine derivative and V-shaped carbazole one (Figure 4). Admixture of small quantities of trifluoroacetic acid (TFA) to the reaction mixture resulted in a significant increase of the reaction yield with almost 100% selectivity towards phenazine derivatives.

![Figure 4](image2)

**Figure 4.** Preparation of tetraalkylphenothio[2,3-a: 2′,3′-h]phenazine from dialkyl-substituted 2-aminoanthraquinone via oxidation with 2,3-dichloro-5,6-dicyano-1,4-benzoquinone (DDQ). R = (triisopropylsilyl)ethynyl [63].

An interesting procedure of the preparation of phenazine derivatives was proposed by Takeda et al. [64]. Using different oxidizing agents 1,1′-binaphthalene-2,2′-diamine was oxidized to two products: U-shaped phenazine derivative and diaza[5]helicene (see Figure 5). The highest selectivity towards dibenzo[a,j]phenazine (77%) was obtained when 1,3-diiodo-5,5-dimethyl-2,4-imidazolidinedione (DIH) was used as an oxidant.
Figure 5. Synthesis of dibenzo[a,j]phenazine and diaza[5]helicene by oxidation of 1′1-binaphthalene-2,2′-diamine [64].

Other, possible methods of azaacenes preparation should also be mentioned. This involves, among others, base-catalyzed condensation of anilines with nitrobenzenes (Wohl-Aue reaction, see Figure 6a) [65] and dehydrative condensation of benzo[1,2,5]oxadiazole-1-oxide with various nucleophiles (Beirut reaction, Figure 6b) [66].

Figure 6. Synthesis of azaacenes via (a) Wohl-Aue and (b) Beirut reactions.

The above discussed reactions have to be carried out at elevated temperatures, which severely limits their use in the syntheses of less thermally stable azaacenes derivatives.

3. Spectroscopic and Electrochemical Properties of Linear Azaacenes

UV-vis-NIR spectra of azaacenes are rich. Strong bands in the UV or visible part of the spectrum are accompanied by a less intensive band in the visible or near infrared part of the spectrum. The latter has a strongly vibronic character with the 0–0 transition being the most intensive (see Figure 7). This least energetic band undergoes increasing bathochromic shift with increasing molecular mass of azaacene, i.e., with number of aromatic rings in the molecule.

Figure 7. UV-vis spectrum of 5,10,15,18-tetraoctyloxy-8,17-dihyronaphtho[2,3-a:2',3'-i]phenazine [67].

As evidenced by cyclic voltammetry, azaacenes undergo either one-step or two-step reversible or quasi-reversible reduction at relatively high potentials (vs. Fc/Fc⁺). This is associated with their high EA value. In the two-step reduction process radical anions are formed in the first step which are then transformed to spinless dianion in the second one. In Figure 8 these processes are depicted using disubstituted tetraazapentacene (quinoxalino[2,3-λ]phenazine) as an example.
Radical anions generation can be evidenced in an EPR spectroelectrochemical experiment or through generation of radical anions directly in the EPR tube, using an appropriate chemical reducing agent [68]. Chemical or electrochemical reduction of azaacenes result in the formation of new UV-vis-NIR bands which can be considered as diagnostic of either the radical anion or spin-less dianion states. By combining UV-vis-NIR and EPR spectroelectrochemistry with cyclic voltammetry it is therefore possible to unequivocally attribute these bands to a given reduction state. To the contrary, oxidation of azaacenes is irreversible and starts at rather high potentials, reflecting their higher IP as compared to acenes.

As already stated, the first reduction potential of azaacenes can be significantly increased and the band gap narrowed, as a result of two main factors or a combination of them: (i) increased number of aromatic rings and (ii) functionalization with electron withdrawing substituents. These effects were discussed in detail in reference [69]. It is instructive to follow the changes of the optical and redox properties in two series of azaacenes depicted in Figure 9. These diazaacenes in series 1 and 2 differed in the number of aromatic rings. Their terminal rings were substituted with two cyano groups. Compounds of series 2 were systematically longer by one ring. The effect of the cyano groups on the LUMO level lowering was evidenced by a significant shift of their first reduction formal potential ($E_{1/2}$) as compared to their non-cyanated analogues [70]. In particular $E_{1/2}$ of 1a was shifted by 600 mV to $-1.1$ V with respect to the same potential of its non-cyanated analogue ($-1.7$ V). Very similar effects were observed for 1b, 1c, 2a and 2b. This implied significantly increased EA values. They are listed in Table 2 together with $E_{1/2}$ and optical band gaps determined from the positions of the 0–0 transitions.
Figure 9. Two series of linear azaacenes synthesized with the goal to investigate the effect of number of rings and electron accepting substituents on their optical and electrochemical properties [69]. R = (triisopropylsilyl)ethynyl group.

Table 2. Electrochemical and optical parameters of series 1 and series 2 diazaacenes [69].

| Compounds | $E_{1/2\text{red}}$ vs. Fc/Fc [V] | EA * [eV] | $E_{\text{opt}}$ [eV] |
|-----------|---------------------------------|-----------|------------------------|
| 1a        | −1.1                            | 3.53      | 2.38                   |
| 1b        | −0.8                            | 3.89      | 1.84                   |
| 1c        | −0.7                            | 4.00      | 1.49                   |
| 2a        | −1.1                            | 3.53      | 2.20                   |
| 2b        | −0.8                            | 3.89      | 1.69                   |
| 2c        | −0.48                           | 4.26      | 1.35                   |

* EA calculated following the procedure proposed in [22].

The effect of the increase in the number of rings is also evident by comparing $E_{1/2}$, EA and $E_{\text{opt}}$ values within one series. For example, EA of 1c increased by 0.47 eV with respect to that measured for 1a whereas the band gap decreased to 1.49 eV, i.e., by 0.83 eV. 1c, 2b and 2c can be considered as low band gap, easily reducible semiconductors. Such materials are highly desirable in organic electronics and photovoltaics.

Halogens are also popular terminal ring substituents lowering the band gap of azaacenes and facilitating their reduction. In Figure 10 two series of compounds are depicted namely di- and tetraazaacenes differently halogenated in their terminal ring. Their spectroscopic parameters are collected in Table 3.

Figure 10. Selected halogen derivatives of di- and tetraazaacenes [50,71]. R = (triisopropylsilyl)ethynyl group.
Table 3. Spectroscopic properties of di- and tetraazaacenes halogenated in their terminal ring, registered for hexane solution (series 3 and 4) [50] and dichloromethane solutions (series 5) [71].

| Compound | \( \lambda_{\text{max abs}} \) [nm] | \( \lambda_{\text{max emi}} \) [nm] | Stokes Shift [nm] |
|----------|---------------------------------|---------------------------------|-----------------|
| 3a       | 572                             | 577                             | 5               |
| 3b       | 602                             | 616                             | 14              |
| 3c       | 620                             | 627                             | 7               |
| 4a       |                                  | 548                             |                 |
| 4b       |                                  | 574                             |                 |
| 4c       |                                  | 598                             |                 |
| 5a       |                                  | 586                             | 4               |
| 5b       |                                  | 602                             | 44              |
| 5c       |                                  | 611                             | 42              |

These data indicate that the introduction of halogen substituents induces a bathochromic shift of the 0–0 transition. This effect is more pronounced in the emission spectra since substitution with halogens leads to increasing Stokes shift, especially in the case of heavier substituents such as Br and I [71].

The reduction potential of halogen-free diazatetracene (3a) is \(-1.2\) V vs. Fc/Fc\(^+\) [50]. Introduction of halogen substituents to its terminal ring results in an increase of this potential in the case of 3b and 3c. Tetraazaacenes 4a, 4b and 4c undergo a two-step reduction with a significant shift of the first reduction potential to \(-0.7\) V and \(-0.6\) V for 4a and 4b, respectively [50]. Slightly smaller shift is observed in the case of 4c.

Electrochemical and spectroscopic properties of longer azaacenes such as tetraazapentacenes [72] or tetraazahexacenes [51] are also very sensitive towards halogen substituents [51,72]. In the case of tetrachloro derivative of bis((tripropan-2-ylsilyl)-ethynyl)-tetrachloroquinolinalino[2,3-b]phenazine significant differences in the absorption spectra of its two isomers are observed. 6a (Cl at 2,3,9,10 positions) absorbs at shorter wavelengths as compared to 6b (Cl at 1,4,8,11 position) (see Table 4). Exchange of chlorine in 6a for bromine to yield 6c also induces a bathochromic shift of the absorption band. Halogen derivatives of tetraazahexacenes (7b and 7c see Figure 11) absorb at near infrared part of the spectrum, concomitantly showing high reduction potential and, by consequence, high electron affinity (Table 4).

Table 4. Spectroscopic and electrochemical data of selected halogen derivatives of tetraazapentacenes [72], tetraazahexacenes [51] and tetraazapentacene with nitro substituent [43].

| Compound | \( E_{\text{g opt}} \) [eV] | \( E_{\text{red vs. Fc/Fc}} \) [V] | EA * [eV] |
|----------|-----------------------------|---------------------------------|-----------|
| 6a       | 737/1.68                    | \(-0.60\)                      | 4.12      |
| 6b       | 752/1.65                    | \(-0.60\)                      | 4.12      |
| 6c       | 744/1.67                    | \(-0.70\)                      | 4.00      |
| 7a       | 862/1.44                    | \(-0.55\)                      | 4.18      |
| 7b       | 1039/1.19                   | \(-0.48\)                      | 4.26      |
| 7c       | 1000/1.24                   | \(-0.53\)                      | 4.20      |
| 8a       | 780/1.59 **                 | \(-0.52\)                      | 4.22      |
| 8b       | 867/1.43 **                 | \(-0.27\)                      | 4.51      |

* EA calculated by following the procedure proposed in [22]. ** \( E_{\text{opt}} \) optical band gaps were calculated from the UV-vis-NIR spectra by determining the wavelength of the absorption onset and converting it from nm to eV.
Figure 11. Selected halogen derivatives of tetraazapentacenes [72] and tetraazahexacenes [51]. R = (triisopropylsilyl)ethynyl group.

Semiconductors of extremely high EA value can be obtained by introducing nitro groups to the terminal ring of tetraazapentacenes (Figure 12 and Table 4).

Figure 12. Selected derivatives of tetraazapentacenes [43]. R = (triisopropylsilyl)ethynyl group.

Azaacenes with electron donating substituents behave differently. They can be treated as donor-acceptor compounds of DA or DAD type showing in selected cases ambipolarity [73]. Introduction of a donor group results in significant lowering of the ionization potential (IP), thus these derivatives are more readily oxidizable than nonsubstituted azaacenes. A series of DAD compounds consisting of two triphenylamine substituents and diazaacene central group can be considered here as an instructive example [73] (see Figure 13, Table 5).

Figure 13. DAD compounds consisting of diazaacene central unit of varying number of aromatic rings and two triphenylamine substituents [73].
Table 5. Spectroscopic and electrochemical properties of azaacenes of increasing number of aromatic rings containing triphenylamine substituents [73]. $E_{\text{ox}}$ vs. Fc/Fc$^+$.  

| Compounds | $\lambda_{\text{max abs}}$ [nm] | $E_{g\text{opt}}$ [eV] | $E_{\text{ox1}}$ [V] | $E_{\text{ox2}}$ [V] |
|-----------|-------------------------------|-----------------|-----------------|-----------------|
| 9a        | 413                           | 2.60            | 0.26            | 0.48            |
| 9b        | 493                           | 2.17            | 0.24            | 0.42            |
| 9c        | 455                           | 2.36            | 0.25            | 0.44            |
| 9d        | 560                           | 1.88            | 0.22            | 0.37            |

UV-vis spectra of these compounds are characterized by a distinct band of charge transfer (CT) character which is being bathochromically shifted with increasing number of aromatic rings in the central azaacene unit. The same trend is observed in their emission spectra. These derivatives undergo a two-step oxidation at relatively high potentials. These two oxidations are related to consecutive removal of two electrons from two triphenylamine substituents. The potential of the first oxidation is only weakly dependent on the number of aromatic rings in the central azaacene unit. However, more pronounced changes are observed in the difference between the first and the second oxidation potentials ($E_{\text{ox2}}-E_{\text{ox1}}$) (Table 5). Decreasing ($E_{\text{ox2}}-E_{\text{ox1}}$) value in the case of azaacenes with larger central units can be rationalized by the fact that larger conjugated cores of the central unit better promote delocalization of the positive charge imposed upon the removal of the first electron, facilitating in this manner the removal of an additional electron from the second triphenylamine substituent.

4. Supramolecular Organization of Linear Azaacenes

Azaacenes, similarly to acenes, readily crystalize and their supramolecular organization is, to a large extent, determined by $\pi-\pi$ stacking of flat molecules. Individual molecules separated by a distance from 0.335 nm to 0.350 nm form slipped studs [69]. Substituents may significantly affect this structural arrangement depending on their size and position in the molecule [69]. Weak noncovalent intermolecular interactions induce sometimes structural dimerization within individual stacking pattern, as in the case of a derivative of phenazine $1a$. In this structural arrangement two intermolecular distances between molecules are observed within the stand. Similar stacking pattern was also observed for diazatetracene containing substituents in its terminal ring ($2a$, [69]). To the contrary, only one intermolecular distance of 0.343 nm within a stack was reported for a different isomer of $1a$, i.e., $1b$ [69].

An interesting example of structural engineering of substituted azaacenes can be found in reference [74]. Introduction of long, branched alkoxy groups to 5,6,13,14-tetraazapentacene resulted in the formation of a smectic-type liquid crystalline phase, stable over a wide temperature range. Liquid crystallinity is advantageous in view of azaacenes application in organic electronics since it facilitates deposition of thin films of desired orientation.

Supramolecular organization of linear azaacenes can be very diversified. These few examples are only aimed at demonstrating typical relationships between the molecular and supramolecular structures of these compounds.

5. Degradation of Linear Azaacenes

Azaacenes are more resistant against photolytic and thermal degradations in the presence of oxygen than the corresponding acenes. This is associated with their significantly higher IP values. Detailed studies of the degradation of naphtho[2,3-$g$]quinoxalines and pyrazino[2,3-$b$]phenazines showed that the degradation starts by oxidation of three rings to their peroxide form and then their further transformation to quinones [75]. The terminal ring turned out to be the most resistant in this process whereas the degradation process started by oxidation of the ring adjacent to it.
Degradation in oxygen-free conditions was significantly slower and proceeded through the formation intermolecular covalent bonds resulting in dimers partially losing their aromatic character [75].

6. Azaacenes Containing Pyrene-Type Units

As stated above, linear azaacenes of higher molecular mass and low N/C may undergo degradative oxidation although they are more resistant against this process than the corresponding acenes. Oxidative degradation occurs less readily in non-linear and more isotropic in shape azaacenes. This can be rationalized through Clar’s aromatic sextet rule. According to it, those polyaromatic compounds are more aromatic and, by consequence, more stable in which more \( \pi \)-sextets (benzoid-type rings) can be identified. A comparison of tetracene with triphenylene can be instructive here. In the former only one Clar’s sextet can be distinguished whereas in the latter—three sextets, imposing its higher stability (see Figure 14).

![Figure 14. Clar’s sextets in triphenylene and tetracene.](image)

From this point of view azaacenes containing fused pyrene-type units seem very promising. As in the case of linear azaacenes, their properties can be tuned by changing the number of aromatic rings, N/C ratio or by introducing electron accepting/electron donating substituents. Structural formulae of selected azaacenes of this type are presented in Figures 15 and 16 whereas their spectroscopic and electrochemical data are collected in Table 6.

| Compounds | \( \lambda_{\text{max, abs}} \) [nm] | \( E_{\text{g, opt}} \) [eV] | \( E_{\text{red vs. Fe/Fc}} \) [$\text{Fc/Fc}^+$] [V] | EA *** [eV] | Compounds | \( \lambda_{\text{max, abs}} \) [nm] | \( E_{\text{g, opt}} \) [eV] | \( E_{\text{red vs. Fe/Fc}} \) [$\text{Fc/Fc}^+$] [V] | EA *** [eV] |
|-----------|-----------------|-----------------|-----------------|---------|-----------|-----------------|-----------------|-----------------|---------|
| 10        | 417             | 2.81            | −0.79           | 3.90    | 19        | 485             | 2.42            | −1.51 *         | 3.05    |
| 11        | 435             | 2.76            | −0.76           | 3.93    | 20        | 441             | 2.51            | −1.69 **        | 2.84    |
| 12        | 440             | 2.70            | −0.73           | 3.97    | 21        | 471             | 2.37            | −1.54 **        | 3.01    |
| 13        | 482             | 2.37            | −0.21           | 4.58    | 22        | 464             | 2.41            | −1.62 **        | 2.92    |
| 14        | 455             | 2.09            | −1.42 *         | 3.15    | 23a       | 443             | 2.65            | −1.68 **        | 2.85    |
| 15        | 571             | 1.85            | −0.71 *         | 3.99    | 23b       | 500             | 2.34            | −1.22 **        | 3.39    |
| 16        | 421             | 2.88            | −1.73 *         | 2.79    | 23c       | 525             | 2.25            | −1.19 **        | 3.43    |
| 17        | 436             | 2.69            | −1.57 *         | 2.98    | 23d       | 529             | 2.22            | −1.16 **        | 3.46    |
| 18        | 425             | 2.78            | −1.44 *         | 3.13    | 23e       | 538             | 2.18            | −1.14 **        | 3.48    |

* recalculated from original data in which potential was given vs. SCE, ** recalculated from original data in which potential was given vs. Ag/AgCl, *** EA calculated following the procedure proposed in [22].
Figure 15. Selected azaacenes containing one pyrene-type units: 10 [76], 11 [76], 12 [76], 13 [76], 14 [77], 15 [77], 16 [76], 17 [78], 18 [78], 19 [78], 20 [79], 21 [79], 22 [79]. R—(tripropyl-2-ylsilyl)ethynyl group introduced to facilitate the deposition of these organic semiconductors on silicon substrates.
Trends observed for this family of azaacenes are similar to those reported for linear azaacenes. The effect of chemical constitution on their electrochemical properties can be briefly summarized as follows: (i) they undergo one or two or even multistep quasi-reversible reduction, the number of redox couples depending on the N/C ratio; (ii) their first reduction potential rises with increasing number of aromatic rings as well as with increasing N/C ratio; (iii) introduction of electron accepting groups to the terminal ring results in the same effect; (iv) although the first reduction potential ($E_{\text{red1}}$) and by consequence electron affinity (EA) almost strictly follow this trend, correlation between $E_{\text{red1}}$ and the optical band gap ($E_{\text{g opt}}$) is not always evident (see Table 6). This means that electron accepting substituents may affect the positions of the LUMO and HOMO levels in a similar manner showing no significant narrowing of the band gap.

The above outlined trends can be illustrated by comparing the first reduction potentials of 16, 11 and 12. These three azaacenes consist of the same core. 16 contains no electron accepting substituents whereas in the remaining two (11 and, 12) the terminal ring was functionalized with different electron withdrawing groups (Figure 15). Introduction of these groups significantly rises the reduction potential, the observed shift depending on the substituent capability of lowering the electron density in the core. Two-fold increase of the number of nitrogen atoms while retaining the same number of aromatic rings and the same substituents results in an increase of the first reduction potential ($E_{\text{red1}}$) by ca. 550 mV (compare the case of 11 and 13). The same trend is observed for pyridazine derivatives (20, 21, 22). Their $E_{\text{red1}}$ increases with increasing electron accepting character of the substituent: phenyl < thienyl < pyridyl.

Figure 16. Azaacenes containing increasing number of pyrene-type units [80].
Azaacenes discussed so far contained one pyrene-type unit. There exist, however, reports on azaacenes comprising two or even more such units \[80\]. In Figure 16 azaacenes of increasing length and increasing number of pyrene units are presented, starting from tetraazahexacene with one pyrene unit and ending up with dodecaazahexadecacene containing three pyrene units \((23a–e)\). One of the characteristic features of this group of compounds is a rather weak dependence of their first reduction potential on the N/C ratio and the number of aromatic rings, observed for azaacenes longer than tetraazahexacene \((23a)\). In fact, extension of the molecule length from octazaoctacene \((23b)\) to dodecaaza-hexadecacene \((23e)\) results in an increase of \(E_{\text{red1}}\) by 60 mV, only. The number of redox couples in compounds \(23a, 23b, 23c, 23d\) and \(23e\) is equal to the number of azaacene units separated by pyrene units.

As already stated, in the discussed compounds quinoxaline and pyrazinoquinoxaline segments are separated by pyrene segments. Their UV-vis-NIR spectra of strongly vibronic nature are typical of azaacenes with some contribution from pyrene. The band ascribed to 0-0 transition of the shortest compound \((23a)\) is hypsochromically shifted with respect to the corresponding bands of the four other compounds by 60 to 90 nm.

### 7. Spatially Extended Azaacenes

Star-shaped azaacenes are typical examples of this group of compounds. In Figure 17 star-shaped azaacenes containing three arms of equal or non-equal length are depicted.

![Figure 17. Chemical structures of star-shaped azaacenes. R = p-decyloxyphenyl group.](image)

An attractive feature of this type of azaacenes is the possibility of individual arms engineering \[81,82\]. Transformation of one arm from quinoxaline-type into pyrazinoquinoxaline results in a significant increase of the first reduction potential and a bathochromic shift of the band ascribed to the 0-0 transition by nearly 40 nm with concomitant increase of \(E_{\text{red1}}\) from \(-1.01\) V to \(-0.71\) V (compare \(24a\) and \(24b\) in Table 7) \[81\]. Consecutive extension of the second and the third arm brings further increase of \(E_{\text{red1}}\) and increasing bathochromic shift of the band ascribed to the 0-0 transition (Table 7). Engineering of an individual arm transfers the induced changes to the remaining two arms because the molecule as a whole is aromatic. Finally, functionalizing the terminal ring of quinoxaline-type arms...
with dicarboxyimide groups (see Figure 18) results in an increase of $E_{\text{red}1}$ to $-0.91$ V and simultaneous increase of the band gap to 2.95 eV [83]. This unexpected finding indicates that lowering of the LUMO level as a result of introducing electron-withdrawing groups was overcompensated by even more pronounced lowering of the HOMO level.

Table 7. Spectroscopic and electrochemical data of star-shaped azaacenes 24 [81], 25 [83].

| Compounds | $\lambda_{\text{max abs}}$ [nm] | $\lambda_{\text{max emi}}$ [nm] | $E_{\text{g opt}}$ [eV] | $E_{\text{red vs. Fc/Fc}^+}$ [V] | $E_{\text{A}^*}$ [eV] |
|-----------|-------------------------------|-------------------------------|---------------------|-------------------------------|---------------------|
| 24a       | 471                           | 552                           | 2.43                | -1.01                         | 3.64                |
| 24b       | 508                           | 593                           | 2.23                | -0.71                         | 3.99                |
| 24c       | 530                           | 605                           | 2.13                | -0.60                         | 4.12                |
| 24d       | 543                           | 615                           | 2.08                | -0.53                         | 4.20                |
| 25        | 399                           | 504                           | 2.95                | -0.91                         | 3.90                |

* $E_{\text{A}^*}$ calculated by following the procedure proposed in [22].

Figure 18. Chemical structure of star-shaped 25 [83]. $R =$ dodecane group.

Spatially extended azaacenes containing cyclic, conjugated central unit deserve a special interest [84,85]. In Figure 19 chemical structures of a series of azaacenes of this type, differing in the nature of their azaacene arms are presented: quinoxaline (26a), benzoquinoxaline (26b) naphthalenequinoxaline (26c), pyrazinophenazine (26d). Spectroscopic and electrochemical properties of these compounds were compared to those determined for molecules mimicking their arms (27) [84].

Spectroscopic properties of 26 a, b, c, d compounds were very similar to those of the corresponding 27 a, b, c, d ones, showing only a small bathochromic shift of their absorption and emission bands (ca. 10–15 nm) (see Table 8) [84]. Electrochemical data were somehow surprising. With the exception of 26a and 27a for which an increase of $E_{\text{red}}$ by 350 mV was observed in the star-like compound, in the three remaining cases a small decrease of $E_{\text{red}}$ by 60 to 100 mV was noticed for 26 b, c, d as compared to 27 b, c, d. This is in contrast to common cases showing that the extension of the pi-system in polyconjugated molecules results in an increase of $E_{\text{red}}$. Evidently the central ring does not transmit the conjugation between individual arms.
Figure 19. Azaacenes with benzenacyclooctaphane central units [84].

Table 8. Spectroscopic and electrochemical data of azaacenes containing conjugated cyclic central unit [84].

| Compounds | $\lambda_{\text{max abs}}$ [nm] | $\lambda_{\text{max emi}}$ [nm] | $E_g$ opt [eV] | $E_{\text{red}}$ vs. Fc/Fc$^+$ [V] | EA $^*$ [eV] |
|-----------|-------------------------------|-------------------------------|----------------|-------------------------------|----------------|
| 26a       | 378                           | 422                           | 2.99           | −1.50                         | 3.90            |
| 26b       | 497                           | 508                           | 2.42           | −1.63                         | 3.06            |
| 26c       | 597                           | 604                           | 1.98           | −1.50                         | 2.91            |
| 26d       | 595                           | 607                           | 2.01           | −1.09                         | 3.06            |
| 27a       | 369                           | 427                           | 3.06           | −1.85                         | 3.54            |
| 27b       | 485                           | 497                           | 2.47           | −1.56                         | 2.65            |
| 27c       | 582                           | 590                           | 2.07           | −1.40                         | 2.99            |
| 27d       | 586                           | 599                           | 2.05           | −1.03                         | 3.18            |

$^*$ EA calculated by following the procedure proposed in [22].

Molecules depicted in Figure 20 can be considered as propeller-shaped azaacenes. The propeller arms consist of quinoxaline, benzoquinoxaline and naphthalenequinoxaline units held together through o-phenylene (28 a, b, c) or m-phenylene (29 a, b, c) linkers. Spectroscopic properties of both types of azaacenes are similar. Their 0–0 transition band is strongly dependent on the length of the arm, undergoing a bathochromic shift with increasing number of aromatic rings. The conjugation between particular arms via phenylene linkers is rather weak since the 0–0 transition bands in 28 a, b, and 29 a, b, c were only slightly bathochromically shifted with respect to the corresponding bands of compounds mimicking their arms (see Table 9).
Figure 20. Molecular structure of propeller-shaped azaacenes [85]. R = (triisopropylsilyl)ethynyl group.

Table 9. Spectroscopic and electrochemical data of propeller-shaped azaacenes of 28 and 29 series [85].

| Compounds | λ_{max abs} [nm] | λ_{max emi} [nm] | E_g opt [nm]/[eV] | E_{red vs. Fc/Fc}^+ [V] | EA * [eV] |
|-----------|-----------------|-----------------|------------------|--------------------------|-----------|
| 28a       | 391             | 413             | 409/3.03         | −1.58                    | 2.97      |
| 28b       | 490             | 500             | 504/2.46         | −1.55                    | 3.00      |
| 28c       | 589             | 594             | 603/2.06         | −1.33                    | 3.26      |
| 29a       | 388             | 433             | 411/3.02         | −1.82                    | 2.68      |
| 29b       | 493             | 506             | 510/2.43         | −1.57                    | 2.98      |
| 29c       | 596             | 601             | 610/2.03         | −1.34                    | 3.25      |

* EA calculated by following the procedure proposed in [22].

These spectroscopic findings were corroborated by electrochemical investigations indicating a very modest increase of the first reduction potential in the propeller-shaped compounds as compared to those determined for molecules mimicking their arms (Table 9).

8. Nonlinear Azaacenes

Nonlinearity of azaacene molecules results in an increased number of Clar’s sextets as compared to their linear analogues, leading to improved stability. There are several types of nonlinear azaacenes [86,87]: L-shaped, Z-shaped, U-shaped spatially extended of perylene- or pyrene-type and others. The effect of nonlinearity on spectroscopic properties of azaacenes can be evidenced by comparing L-shaped diazapentacene with its linear analogue and linear diazatetracene (see Figure 21). Their spectroscopic data are collected in Table 10. It is evident that nonlinearity induces a strong hypsochromic shift of the band corresponding to the 0–0 transition by nearly 150 nm. E_{g opt} of 32 is even larger than that of linear diazatetracene—31 [88].

Figure 21. Chemical structures of: 6,11-bis((triisopropylsilyl)ethynyl)benzo[b]phenazine (30) and 6,13-bis-((triisopropylsilyl)ethynyl)naphtho[2,3-b]phenazine (31) and L-shaped 8,13-bis((triisopropylsilyl)ethynyl)dibenzo[a,i]phenazine (32) [88]. R = (triisopropylsilyl)ethynyl group.
Table 10. Spectroscopic data, redox potentials and electron affinities (EA) of linear, L-shaped and Z-shaped azaacenes [88,89].

| Compound | $\lambda_{\text{max abs}}$ [nm] | $\lambda_{\text{max emi}}$ [nm] | $E_{\text{g opt}}$ [nm]/[eV] | $E_{\text{red vs. Fc/Fc}}$ [V] | EA * [eV] |
|----------|-------------------------------|-------------------------------|-----------------------------|---------------------------------|-----------|
| 30       | 570                           | 577                           | 585/2.12                    | $-1.23$ **                      | 3.38      |
| 31       | 693                           | 699                           | 709/1.75                    | $-1.05$ **                      | 3.59      |
| 32       | 544                           | 553                           | 558/2.22                    | $-1.79$ **                      | 2.72      |
| 33       | 681                           | 694                           | $-/1.94$                    | $-0.68$                         | 4.03      |
| 34       | 653                           | 699                           | $-/1.75$                    | $-0.76$                         | 3.93      |
| 35       | 712                           | 688                           | $-/1.75$                    | $-0.51$                         | 4.23      |

* EA calculated by following the procedure proposed in [22] ** $E_{\text{red}}$ calculated from data reported in [88].

Comparison of spectroscopic data obtained for linear 33, L-shaped 34 and Z-shaped 35 azaacenes (Figure 22 and Table 10) shows that the 0–0 transition band of 33 is bathochromically shifted by only 31 nm with respect to the corresponding band in linear diazapentacene 34. Similarly, its $E_{\text{g opt}}$ is only slightly narrower by 0.06 eV as compared to $E_g$ of 34. This finding should be considered as a spectroscopic evidence of a very weak conjugation between tetraazapentacene linear segments of this Z-shaped molecule.

Figure 22. Molecular formulae of linear tetraazapentacene (6,13-bis((triisopropylsilyl)ethynyl)-5,7,12,14-tetraazapentacene) 33, L-shaped tetraazahexacene (8,15-bis((triisopropylsilyl)-ethynyl)-benzo[a]quinoxalino[2,3-i]phenazine) 34 and Z-shaped 6,12,19,25-tetra((triisopropylsilyl)ethynyl)-quinoxalino 2,3-i|quinoxalino [2',3':6',7']quinoxalino[2',3':5,6]nafto[2,3-a]phenazine 35 [89]. R = (triisopropylsilyl)ethynyl group.

Diazaacenes usually exhibit one quasi-reversible redox couple at negative potentials with respect to Fc/Fc+. In Table 10 reduction potentials of L-shaped diazapentacene 32, linear diazapentacene 31 and linear diazatetraacene 30 are compared. The lowest reduction potential is observed for the L-shaped compound, which is even lower than that measured for the linear one 30. This can be considered as a manifestation of less efficient conjugation of 32 due to the presence of two mutually independent Clar’s sextets.

Tetraazaacenes, such as linear tetraazapentacene 33, L-shaped tetraazapentacene 34 as well as Z-shaped octaazaundecacene 35, usually exhibit two redox couples at negative potentials, ascribed to two consecutive one-electron reductions to a radical anion and dianion, respectively. Furthermore, in this case the L-shaped molecule shows lower reduction potential than the linear one, although the latter is shorter by one ring. Significant increase of the first reduction potential is observed for the Z-shaped compound 35. This molecule consists of two linear tetraazapentacene segments connected by an aromatic ring. An increase of the potential by 0.17 V with respect to that of linear tetraazapentacene 33 indicates that the conjugation extends over the connecting (central) aromatic ring.
Solution processable Z-shaped azaacenes which can be considered as derivatives of dinaphtho[2,3-a:2',3'-h]phenazine (see Figure 23) were synthesized by Goto et al. [63] and Kotwica et al. [37,38]. Alkoxy groups introduced to 37 induce solution processability but in addition, due to their electron donating properties, they lower the band gap which is slightly lower than the gap of 36 and significantly lower that the gap of 42 – unsubstituted dibenzophenazine (see Table 11). 38 and 39, i.e., benzo[h]benz[5,6]acridino[2,1,9,8-klnma]acridines were reported by Kotwica et al. [61]. These compounds can be considered as two L-shaped azatetracenes fused together. Note a close similarity of the absorption and emission spectra as well as the reduction potential of 37 and 39. 40 exhibits a large gap and a very low reduction potential. Consistent with these finding it emits ultraviolet radiation.

Figure 23. Azaacenes 36 [63], 37 [60], 38 [90], 39 [61], 40 and 41 [91], 42 [64]. R₁ and R₃ = (triisopropylsilyl)ethynyl, ₅₆ = octyl group.

Table 11. Spectroscopic and electrochemical data of azaacenes 36 [63], 37 [60], 38 [90], 39 [61], 40 and 41 [91], 42 [64].

| Compounds | λ_{max \text{ abs.}} [nm] | λ_{max \text{ emi.}} [nm] | E_{g \text{ opt}} [eV] | E_{\text{red vs. } \text{Fc/Fc}^+} [V] | EA [eV] |
|-----------|----------------|----------------|----------------|----------------|----------|
| 36        | 499           | 501           | -             | -1.20          | 3.41     |
| 37        | 493           | 515           | 2.40          | -1.63          | 2.91     |
| 38        | 554           | -             | 2.23          | -              | -        |
| 39        | 515           | 524           | 2.33          | -1.62          | 2.92     |
| 40        | 369           | 370           | 4.11          | -2.02          | 2.45     |
| 41        | 423           | 431           | 3.18          | -1.77          | 2.74     |
| 42        | 416           | 425           | -             | -1.88          | 2.61     |

In summary, the possibilities of precise tuning of azaacenes properties should be pointed out, which are more numerous as compared to acenes. Electronic, optical and redox properties can be modified not only by increasing the number of aromatic rings but also by changing N/C ratio and the mode of nitrogen atoms distribution within
the molecule. Finally, changing the shape of molecules (linear vs. L-shaped, Z-shaped, U-shape, condensed etc.) not only leads to alteration of their redox properties but also may result in distinctly different supramolecular organizations. Data reported in reference [92] should be considered as an instructive example of such tuning of properties. 6,13-bis((triisopropylsilyl)ethynyl)-1,4-diazapentacene, i.e., diazapentacene containing two nitrogen atoms in its terminal ring shows hole-only electrical transport in the field effect transistor configuration with a mobility of 1.2 cm²V⁻¹s⁻¹. Transferring these two nitrogen atoms to one of the inner rings as in the case of 6,13-bis((triisopropylsilyl)ethynyl)-5,14-diazapentacene transforms this hole transporting semiconductor into an ambipolar one. Once again, in the FET configuration values of 0.22 cm²V⁻¹s⁻¹ and 1.1 cm²V⁻¹s⁻¹ were measured for electron and holes mobilities, respectively. Increasing the number of nitrogen atoms from two to four gives rise to a purely electron transporting (n-type) semiconductor, exhibiting electron mobility of 3.3 cm²V⁻¹s⁻¹ [92]. Specific properties, including processability, can also be tuned by functionalization. Functionalization of dinaphtho[2,3-a′,2′,3′-h]phenazine with alkoxy groups [60] lowers the band gap of this electroluminophore and increases its solubility. Additionally, strong dependence of the melting temperature on the substituent length is observed. In particular, it decreases from nearly 200 °C to below 100 °C when butoxy substituents are replaced by dodecyloxy ones, with redox and spectroscopic properties remaining essentially the same. This appropriately functionalized dinaphtho[2,3-a′,2′,3′-h]phenazine can be conveniently melt-processed.

9. Applications of Azaacenes

High stability of azaacene radical anions and in some cases radical cations facilitates their use as components of organic and hybrid (organic/inorganic) electronic and optoelectronic devices. This is important because in operating conditions organic semiconductors are being temporarily transformed from their neutral form to the ionic one, due to electron or hole injection. Several light emitting diodes of excellent efficiency were fabricated where azaacenes served as electroluminophores in active layers [82,93–95] or as electron transporting or hole blocking layers [96]. An important drawback of classical organic light emitting diodes should be pointed out. In these devices electrons and holes injected from the electrodes form excitons. Only 25% of them are singlet excitons which may undergo radiative recombination generating light. 75% are triplet excitons which do not contribute to the generation of light. Thus, the maximum internal quantum efficiency in classical organic diodes cannot exceed 25%. However, if the energy difference between the singlet and the triplet states is small, the singlet state can be thermally repopulated through the reversed intersystem crossing (RISC), transforming inactive triplet excitons into singlet ones. This phenomenon is called “thermally activated delayed fluorescence” (TADF) and is very intensively studied in recent years, because TADF diodes can reach internal efficiencies approaching 100%. Several azaacenes were reported as efficient TADF emitters in recent years, among them substituted dibenzo[a,c]phenazines 52, 53 (see Table 12) [97] or 10-(acenaphtho[1′,2′:5,6]pyrazino[2,3-f][1,10]phenanthroin-12-yl)-10H-phenoxazine 46 and 7-(acenaphtho[1′,2′:5,6]pyrazino[2,3-f][1,10]phenanthroin-12-yl)-7H-benzol[c]phenoxazine 47 (Table 12) [98]. Table 12 presents chemical formulae of azaacenes used as components of OLEDs together with selected parameters of the fabricated test devices (positions 43–64).

In recent years, in the domain of organic solar cells, significant research effort was directed towards the design and fabrication of “all organic” solar cells in which fullerene derivative-type acceptors, typically used in “bulk heterojunction” devices, were replaced by purely organic acceptors (see for example [99,100]). Various azaacenes were tested as acceptors in such devices [36,37,101]. Formulae of selected azaacene-type acceptors are listed in Table 12 (positions 65–68), together with parameters of the test devices.

In the past decade, devices in which the active layer consists of various derivatives of lead or tin halides of perovskite structure, the so called “perovskite solar cells”, dominated the research on hybrid (inorganic/organic) photovoltaics [102], despite several
problems connected to their stability in operating conditions. However, significant effort was achieved in few recent years in this respect, as far as elucidation of degradation mechanisms and stability improvements are concerned [103]. Nitrogen atoms-rich azaacene derivatives are very well suited to serve as electron transporting layers in perovskite solar cell [104–106]. Chemical formulae of azaacenes used in perovskite-type devices are presented in Table 12 (positions 69–74) together with selected device parameters.

Table 12. Structures of azaacenes and their application in devices.

| Compound | Device’s Properties |
|----------|---------------------|
|          | Emitters in organic light emitting diode |
|          | maximum luminance 1000 cd m$^{-2}$ *  |
|          | efficiency 5 cd A$^{-1}$  |
|          | EQE = 0.53%  |
|          | $\lambda_{\text{max emi}}$ = 540 nm *  |
| 43       | [93]               |
|          | maximum luminance 2321 cd m$^{-2}$  |
|          | efficiency 0.79 cd A$^{-1}$,  |
|          | $\lambda_{\text{max emi}}$ = 547 nm  |
| 44       | [85]               |
|          | maximum luminance 5792 cd m$^{-2}$ *  |
|          | efficiency 2.88 cd A$^{-1}$  |
|          | $\lambda_{\text{max emi}}$ = 502 nm  |
| 45       | [113]              |
|          | maximum luminance > 600 cd m$^{-2}$ *  |
|          | efficiency 5.4 cd A$^{-1}$  |
|          | EQE = 7.2%  |
|          | $\lambda_{\text{max emi}}$ = 640 nm  |
| 46       | [98]               |
|          | maximum luminance > 300 cd m$^{-2}$ *  |
|          | efficiency 0.8 cd A$^{-1}$  |
|          | EQE = 2.0%  |
|          | $\lambda_{\text{max emi}}$ = 665 nm  |
| 47       | [98]               |
|          | maximum luminance > 300 cd m$^{-2}$ *  |
|          | EQE = 22.80%  |
|          | $\lambda_{\text{max emi}}$ = 698 nm  |
| 48       | [114]              |
| Compound | Device's Properties |
|----------|---------------------|
| ![Compound 49](image) | maximum luminance > 10,000 cd m$^{-2}$  
EQE = 26.00%,  
[115] |
| ![Compound 50](image) | efficiency 51.8 cd A$^{-1}$  
EQE = 21.8%,  
$\lambda_{\text{max emi}} = 577$ nm  
[116] |
| ![Compound 51](image) | efficiency 47.1 cd A$^{-1}$  
EQE = 23.8%,  
$\lambda_{\text{max emi}} = 587$ nm  
[116] |
| ![Compound 52](image) | maximum luminance > 6000 cd m$^{-2}$  
EQE = 19.4%,  
$\lambda_{\text{max emi}} = 580$nm  
[97] |
| ![Compound 53](image) | maximum luminance > 10,000 cd m$^{-2}$  
EQE = 22.1%,  
$\lambda_{\text{max emi}} = 547$ nm  
[97] |
| ![Compound 54](image) | maximum luminance > 1000 cd m$^{-2}$  
EQE = 3.3%,  
$\lambda_{\text{max emi}} = 536$ nm phosphorescence  
[33] |
Table 12. Cont.

| Compound | Device’s Properties |
|----------|---------------------|
| ![Compound 55](image55.png) | maximum luminance $8600 \text{ cd m}^{-2}$, EQE = 0.33%, $\lambda_{\text{max emi}} = 511 \text{ nm}$ [117] |
| ![Compound 56](image56.png) | maximum luminance $> 200 \text{ cd m}^{-2}$, efficiency $1 \text{ cd A}^{-1}$, $\lambda_{\text{max emi}} = 590 \text{ nm}$ [70] |
| ![Compound 57](image57.png) | maximum luminance $593 \text{ cd m}^{-2}$ *, efficiency $0.85 \text{ cd A}^{-1}$, $\lambda_{\text{max emi}} = 502 \text{ nm}$ [118] |
| ![Compound 58](image58.png) | maximum luminance $20780 \text{ cd m}^{-2}$, EQE = 3.1%, $\lambda_{\text{max emi}} = 500 \text{ nm}$ [119] |
| ![Compound 59](image59.png) | maximum luminance $> 500 \text{ cd m}^{-2}$ *, efficiency 11.6 cd A$^{-1}$, EQE = 10.4%, $\lambda_{\text{max emi}} = 456 \text{ nm}$ [120] |
| ![Compound 60](image60.png) | maximum luminance $920 \text{ cd m}^{-2}$ *, efficiency 80 cd A$^{-1}$, EQR = 0.53%, $\lambda_{\text{max emi}} = 495 \text{ nm}$ [31] |
Table 12. Cont.

| Compound | Device’s Properties |
|----------|---------------------|
| ![Image](61) | maximum luminance 595 cd m\(^{-2}\)*, efficiency 7.0 cd A\(^{-1}\), EQE = 2.0%, \(\lambda_{\text{max emi}} = 520\) nm* [15] |
| ![Image](62) | maximum luminance 80 cd m\(^{-2}\), \(\lambda_{\text{max emi}} = 508\) nm [89] |
| ![Image](63) | efficiency 2.13 cd A\(^{-1}\), EQE = 4.61%, \(\lambda_{\text{max emi}} = 440\) nm [121] |
| ![Image](64) | EQE = 0.05%, \(\lambda_{\text{max emi}} = 970\) nm [122] |

Acceptors for solar cells

| Compound | Properties |
|----------|------------|
| ![Image](65) | \(V_{\text{OC}} = 0.81\) V, FF = 37.3%, \(J_{\text{SC}} = -6.77\) mA cm\(^{-2}\), PCE = 2.03% [101] |
| ![Image](66) | \(V_{\text{OC}} = 0.75\) V, FF = 38.3%, \(J_{\text{SC}} = -5.66\) mA cm\(^{-2}\), PCE = 1.62% [35] |
### Table 12. Cont.

| Compound | Device's Properties |
|----------|---------------------|
| ![Compound 67](image) | $V_{OC} = 0.80 \text{ V}$, $FF = 41\%$, $J_{SC} = -6.6 \text{ mA cm}^{-2}$, $PCE = 2.19\%$ [37] |
| ![Compound 68](image) | $V_{OC} = 0.85 \text{ V}$, $FF = 36\%$, $J_{SC} = -7.51 \text{ mA cm}^{-2}$, $PCE = 2.28\%$ [36] |

#### Electron Transporting Layers for Perovskite Solar Cells

| ![Compound 69](image) | $\mu_e = 2.8 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, $\mu_e = 2.7 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (100 °C) [104] |
| ![Compound 70](image) | $\mu_e = 4.7 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [105] |
| ![Compound 71](image) | $\mu_e = 1.73 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [106] |
| ![Compound 72](image) | $\mu_e = 1.9 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [123] |
Table 12. Cont.

| Compound | Device’s Properties |
|----------|---------------------|
| ![Compound 73](image1) | $\mu_e = 5.13 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [124] |
| ![Compound 74](image2) | $\mu_e = 1.68 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (doped by Et$_3$N) [125] |
| ![Compound 75](image3) | $\mu_e = 7 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [69] |
| ![Compound 76](image4) | $\mu_e = 7 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [126] |
| ![Compound 77](image5) | $\mu_e = 9 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [127] |
| ![Compound 78](image6) | $\mu_e = 0.14 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, $\mu_h = 0.12 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [109] |

Active layers in organic field effect transistors
Table 12. Cont.

| Compound | Device’s Properties |
|----------|---------------------|
| ![Compound 79](image1.png) | $\mu_e = 0.15 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, $\mu_h = 0.11 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [108] |
| ![Compound 80](image2.png) | $\mu_e = 0.05 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, $\mu_h = 4 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [107] |
| ![Compound 81](image3.png) | $\mu_e = \text{ from } 1 \text{ do } 3.3 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (depend of temperatures of substrate—from $25 \degree \text{C}$ to $100 \degree \text{C}$) $\mu_e = 11 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (measured in vacuum) [107] |
| ![Compound 82](image4.png) | $\mu_e = 3.5 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [129] |
| ![Compound 83](image5.png) | $\mu_e = 27.8 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (measured in vacuum) [111] |
| ![Compound 84](image6.png) | $\mu_e = 0.56 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [130] |
| ![Compound 85](image7.png) | $\mu_e = 3.7 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [131] |
| ![Compound 86](image8.png) | $\mu_h = 0.06 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [110] |
| Compound | Device’s Properties |
|----------|---------------------|
| ![Compound 1](image1.png) | $\mu_h = 8.1 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [132] |
| ![Compound 2](image2.png) | $\mu_e = 8.1 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, $\mu_h = 2 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [133] |
| ![Compound 3](image3.png) | **Self-assembled monolayers (SAM) for functionalization of gold electrodes for organic field effect transistors** |
| ![Compound 4](image4.png) | on/off ratio increased from $10^2$ to $10^4$, $\mu_e$ from $10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ to $10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [134] |
| ![Compound 5](image5.png) | **Photocathode** |
| ![Compound 6](image6.png) | current density 0.13 mA cm$^{-2}$ at $-0.13$ V (Evs RHE) [135] |
| ![Compound 7](image7.png) | Detected [112] |
As already mentioned, electrical transport properties of azaacenes depend on four major factors: N/C ratio, the number of aromatic rings, the presence of electron donating (electron accepting) substituents and finally the shape of the molecule—linear vs. L-shaped, Z-shaped, U-shaped etc. Several azaacenes were used as active components of n-channel field effect transistors (FETs), some of them are shown in Table 12 (80 [107]). Azaacenes of lower N/C can be ambipolar, i.e., transporting either holes or electrons depending on polarity. Some ambipolar azaacenes are depicted in Table 12 (79 [108], 78 [109]). Azaacenes exhibiting only p-type electrical transport and for these reasons used in p-channel FETs were also reported (86 in Table 12) [110]. In Table 12 (positions 75–88) chemical formulae of selected azaacenes are shown tested as components of active layers in field effect transistors together with their charge carriers mobilities measured in the FET configuration. It should be noted that tetracholorotetraazapentacene reported in [111] exhibited extremely high electron mobility of ca. 28 cm² V⁻¹ s⁻¹.

Application of azaacenes as components of organic memory devices of WORM (write-once-read-many-times) or FLASH (storage that can be electrically erased and re-programmed) have drawn a significant research interest in recent years. Test devices described in [29] exhibited improved stability both in neutral states as well as in on states of reduced resistivity. Selected azaacenes used as components of organic memory devices are presented in Table 12 (92–95).

To summarize, azaacenes exhibit several interesting properties difficult to match by other families of electroactive organic compounds. Their redox, luminescent and electrical transport properties can be precisely tuned by changing the number of aromatic rings being in conjugation, N/C ratio and nitrogen atoms distribution. Further modifications are possible by introducing electron accepting or electron donating substituents. Their solution
or melt processability can be induced or improved by attaching appropriate side groups which, if properly designed, do not worsen their precious electrical transport properties. Thus, it is clearly evident that further development of azaacene-based materials has to be inherently associated with concomitant progress in the synthesis of new functional azaacene derivatives with functions intentionally designed for a given application. This is a very promising direction, taking into account quick progress in heterocyclic chemistry with elaboration of new catalytic systems leading to significantly simplified reaction pathways and resulting in improved yields. One direction in the design of new azaacenes seems extremely interesting, although, so far, little explored. This is self-assembly through molecular recognition, i.e., bio-inspired approach. Introducing substituents capable of forming multiple, parallel intermolecular hydrogen bonds should facilitate formation of ordered monolayers on appropriate substrates such as HOPG or gold. Moreover, appropriate bio-inspired functionalization may facilitate controlled layer by layer deposition of these semiconductors, governed by molecular recognition [139]. Nanoprocessing of this type is of crucial importance for further development of organic nanodevices.

10. Conclusions

This short review demonstrates that azaacenes are extremely interesting compounds for electronic and optoelectronic applications. They are in general more stable and more resistant to degradative oxidation than the corresponding acenes. Moreover, by changing their N/C ratio as well as distribution mode of N-atoms in the molecule their optical, electronic and redox properties can be tuned in a wide range. Additional modification of their properties can be achieved by changing the molecular shape (linear vs. L-shaped, Z-shaped, U-shaped, fused, etc.) and by introduction of substituent of electron accepting or electron donating character. Finally, they can be rendered solution and melt processable by attaching appropriate flexible solubilizing groups to the conjugated core or substituents lowering the melting temperature. For all these reasons various types of azaacenes are studied as active components of field effect transistors, phototransistors, light emitting diodes, memory devices and others.

Author Contributions: K.K. literature search and writing; I.W. literature search and A.P. writing. All authors have read and agreed to the published version of the manuscript.

Funding: This work was supported by the National Science Centre of Poland, Grant No. 2019/32/C/ST5/00179 and Grant No. 2019/33/B/ST5/00582.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Conflicts of Interest: The authors declare no conflict of interest.

References
1. Bunz, U.H.F. The larger linear N-heteroacenes. Acc. Chem. Res. 2015, 48, 1676–1686. [CrossRef] [PubMed]
2. Pron, A.; Gawrys, P.; Zagorska, M.; Djurado, D.; Demadrille, R. Electroactive materials for organic electronics: Preparation strategies, structural aspects and characterization techniques. Chem. Soc. Rev. 2010, 39, 2577–2632. [CrossRef] [PubMed]
3. Xie, W.; Li, B.; Cai, X.; Li, M.; Qiao, Z.; Tang, X.; Liu, K.; Gu, C.; Ma, Y.; Su, S.-J. Thiophene disubstituted benzothiadiazole derivatives: An effective planarization strategy toward deep-red to near-infrared (NIR) organic light-emitting diodes. Front. Chem. 2019, 7, 276. [CrossRef] [PubMed]
4. Ahrens, L.; Hahn, S.; Rominger, F.; Freudenberg, J.; Bunz, U.H.F. N-Acenoacenes. Chem. Eur. J. 2019, 25, 14522–14526. [CrossRef]
5. Takeya, J.; Yamagishi, M.; Tominari, Y.; Hirahara, R.; Nakazawa, Y.; Nishikawa, T.; Kawase, T.; Shimoda, T.; Ogawa, S. Very high-mobility organic single-crystal transistors with in-crystal conduction channels. Appl. Phys. Lett. 2007, 90, 102120. [CrossRef]
6. Ono, K.; Totani, H.; Hiei, T.; Yoshino, A.; Saito, K.; Eguchi, K.; Tomura, M.; Nishida, J.; Yamashita, Y. Photooxidation and reproduction of pentacene derivatives substituted by aromatic groups. Tetrahedron 2007, 63, 9699–9704. [CrossRef]
7. Mallo, G.; Mulas, G.; Cappellini, G.; Joblin, C. Time-dependent density functional study of the electronic spectra of oligoacenes in the charge states $-1$, $0$, $+1$, and $+2$. Chem. Phys. 2007, 340, 43–58. [CrossRef]
35. Ahrens, L.; Butscher, J.; Brosius, V.; Rominger, F.; Freudenberg, J.; Vaynzof, Y.; Bunz, U.H.F. Azaacene dimers: Acceptor materials with a twist. Chem. Eur. J. 2020, 26, 412–418. [CrossRef] [PubMed]

36. Lami, V.; Leibold, D.; Fassl, P.; Hofstetter, Y.J.; Becker-Koch, D.; Bieger, P.; Paulus, F.; Hopkinson, P.E.; Adams, M.; Bunz, U.H.F.; et al. N-Heteroacenes as a new class of non-fullerene electron acceptors for organic bulk-heterojunction photovoltaic devices. Sol. RRL 2017, 1, 1700053. [CrossRef]

37. Leibold, D.; Lami, V.; Hofstetter, Y.J.; Becker-Koch, D.; Weu, A.; Bieger, P.; Paulus, F.; Bunz, U.H.F.; Hopkinson, P.E.; Bakulin, A.A.; et al. Triptycynel-phenazino-thiadiazole as acceptor in organic bulk-heterojunction solar cells. Org. Electron. 2018, 57, 285–291. [CrossRef]

38. Fischer, O.; Hepp, E. Oxidation des orthophenylendiamins. Ber. Dtsch. Chem. Ges. 1889, 22, 355–359. [CrossRef]

39. Brosius, V.; Müller, M.; Borstelmann, J.; Rominger, F.; Freudenberg, J.; Bunz, U.H.F. Azaacenodibenzosuberones. [CrossRef]

40. Ahrens, L.; Tverskoy, O.; Weigold, S.; Ganschow, M.; Rominger, F.; Freudenberg, J.; Bunz, U.H.F. (Aza)pentacenes clipped into a ring: Stabilization of large (aza)acenes. Angew. Chem. Int. Ed. 2021, 60, 9270–9273. [CrossRef]

41. Zhang, Z.; Zhang, Q. Recent progress in well-defined higher azaacenes (n ≥ 6): Synthesis, molecular packing, and applications. Mater. Chem. Front. 2020, 4, 3419–3432. [CrossRef]

42. Weng, S.-Z.; Shukla, P.; Kuo, M.-Y.; Chang, Y.-C.; Sheu, H.-S.; Chao, I.; Tao, Y.-T. Diazapentacene derivatives as thin-film transistor materials: Morphology control in realizing high-field-effect mobility. ACS Appl. Mater. Interfaces 2009, 1, 2071–2079. [CrossRef]

43. Tverskoy, O.; Rominger, F.; Peters, A.; Himmel, H.-J.; Bunz, U.H.F. An efficient synthesis of tetraazapentacenes. Angew. Chem. Int. Ed. 2011, 50, 3557–3560. [CrossRef]

44. Tong, C.; Zhao, W.; Luo, J.; Mao, H.; Chen, W.; Chan, H.S.O.; Chi, C. Large-Size linear and star-shaped dihydropyrazine fused pyrazinacenes. Org. Lett. 2012, 14, 494–497. [CrossRef]

45. Seillan, C.; Brisset, H.; Siri, O. Efficient synthesis of substituted dihydrotetraazapentacenes. Org. Lett. 2008, 10, 4013–4016. [CrossRef] [PubMed]

46. Liu, T.-C.; Lin, K.-C.; Kon-no, M.; Lee, C.-C.; Shinmyozu, T. Quinoxaline-embedded polyacenoquinone esters: Synthesis, electronic properties, and crystal structure. Org. Lett. 2011, 13, 4588–4591. [CrossRef] [PubMed]

47. Brosius, V.; Müller, M.; Borstelmann, J.; Rominger, F.; Freudenberg, J.; Bunz, U.H.F. Azaacenodibenzoquinones. J. Org. Chem. 2020, 85, 296–300. [CrossRef]

48. Zhu, G.; Zhang, G. Access to a phthalazine derivative through an angular cis-quinacridone. J. Org. Chem. 2021, 86, 1198–1203. [CrossRef] [PubMed]

49. Gevry, P.; Marszalek, T.; Barltin, E.; Kucinska, M.; Ulanski, J.; Zagorska, M. Novel, low-cost, highly soluble n-type semiconductors: Tetraazaanthracene tetraesters. Org. Lett. 2011, 13, 6090–6093. [CrossRef] [PubMed]

50. Lindner, B.D.; Engelhart, J.U.; Märken, M.; Tverskoy, O.; Appleton, A.L.; Rominger, F.; Hardcastle, K.I.; Enders, M.; Bunz, U.H.F. Synthesis and optical properties of diaza-and tetraazaatetracenes. Chem. Eur. J. 2012, 18, 4627–4633. [CrossRef]

51. Engelhart, J.U.; Lindner, B.D.; Schaffroth, M.; Schrempp, D.; Tverskoy, O.; Bunz, U.H.F. Substituted tetraaza- and hexaaazaacenes and their N,N′-dihydro derivatives: Syntheses, properties, and structures. Chem. Eur. J. 2015, 21, 8121–8129. [CrossRef]

52. Bunz, U.H.F.; Engelhart, J.U. The palladium way to N-heteroacenes. Chem. Eur. J. 2016, 22, 4680–4689. [CrossRef]

53. Schwaben, J.; Münster, N.; Klues, M.; Breuer, T.; Hofmann, P.; Harms, K.; Witte, G.; Koert, U. Efficient syntheses of novel fluoro-substituted pentacenes and azapentacenes: Molecular and solid-state properties. Chem. Eur. J. 2015, 21, 13758–13771. [CrossRef]

54. Zhang, X.-X.; Buchwald, S.L. Efficient synthesis of N-aryl-aza-crown ethers via palladium-catalyzed amination. J. Org. Chem. 2000, 65, 8027–8031. [CrossRef]

55. Lindner, B.D.; Engelhart, J.U.; Tverskoy, O.; Appleton, A.L.; Rominger, F.; Peters, A.; Himmel, H.-J.; Bunz, U.H.F. Stable hexacenes through nitrogen substitution. Angew. Chem. Int. Ed. 2011, 50, 8588–8591. [CrossRef]

56. He, Z.; Mao, R.; Liu, D.; Miao, Q. Highly electron-deficient hexaazapentacenes and their dihydro precursors. Org. Lett. 2012, 14, 4190–4193. [CrossRef] [PubMed]

57. China Raju, B.; Veera Prasad, K.; Saidachary, G.; Sridhar, B. A novel approach for C–C, C–N, and C–O bond formation reactions: A facile synthesis of benzoazepine, quinoxaline, and phenoxy derivatives via ring opening of benzoxepines. Org. Lett. 2014, 16, 420–423. [CrossRef]

58. Sikari, R.; Chakraborty, G.; Guin, A.K.; Paul, N.D. Nickel-catalyzed [4 + 2] annulation of nitriles and benzylamines by C–H/N–H activation. J. Org. Chem. 2021, 86, 279–290. [CrossRef] [PubMed]

59. Kotwica, K.; Bujak, P.; Wamal, D.; Materna, M.; Skoroka, L.; Gunka, P.A.; Nowakowski, R.; Golec, B.; Lussczynska, B.; Zagorska, M.; et al. Indanthyone dyes revisited after sixty years. Chem. Commun. 2014, 50, 11543–11546. [CrossRef]

60. Kotwica, K.; Bujak, P.; Wamal, D.; Pieczonka, A.; Wiosna-Salyga, G.; Gunka, P.A.; Jaroch, T.; Nowakowski, R.; Lussczynska, B.; Witkowska, E.; et al. Structural, spectroscopic, electrochemical, and electroluminescent properties of tetraalkoxydinaphthozenes: New solution-processable nonlinear azaacenes. J. Phys. Chem. C 2015, 119, 10700–10708. [CrossRef] [PubMed]

61. Kotwica, K.; Bujak, P.; Data, P.; Krzywiec, W.; Wamal, D.; Gunka, P.A.; Skoroka, L.; Jaroch, T.; Nowakowski, R.; Pron, A.; et al. Soluble flavanthrone derivatives: Synthesis, characterization, and application to organic light-emitting diodes. Chem. Eur. J. 2016, 22, 7978–7986. [CrossRef] [PubMed]
90. Zhang, L.; Fonari, A.; Zhang, Y.; Zhao, G.; Coropeanu, V.; Hu, W.; Parkin, S.; Brédas, J.-L.; Briseno, A.L. Triisopropylsilyl-ethynyl-functionalized graphene-like fragment semiconductors: Synthesis, crystal packing, and density functional theory calculations. Chem. Eur. J. 2013, 19, 17907–17916. [CrossRef] [PubMed]

91. Schneider, J.A.; Perepichka, D.F. A new approach to polyyclic azaarenes: Visible-light photolysis of vinyl azides in the synthesis of diazabenzopyrene and diazaperylene. J. Mater. Chem. C 2016, 4, 7269–7276. [CrossRef]

92. Liang, Z.; Tang, Q.; Mao, R.; Liu, D.; Xu, J.; Miao, Q. The position of nitrogen in N-heteropentacenes matters. Adv. Mater. 2011, 23, 5514–5518. [CrossRef]

93. Lindner, B.D.; Zhang, Y.; Höfle, S.; Berger, N.; Teusch, C.; Jesper, M.; Hardcastle, K.I.; Qian, X.; Lemmer, U.; Colsmann, A.; et al. N-fused quinoxalines and benzoquinolines as attractive emitters for organic light emitting diodes. J. Mater. Chem. C 2013, 1, 5718–5724. [CrossRef]

94. Li, J.; Yan, F.; Gao, J.; Li, P.; Xiong, W.-W.; Zhao, Y.; Sun, X.W.; Zhang, Q. Synthesis, physical properties and OLED performance of azatetrazenes. Dyes Pigm. 2015, 112, 93–98. [CrossRef]

95. Gu, P.-Y.; Wang, N.; Wu, A.; Wang, Z.; Tian, M.; Sun, X.W.; Zhang, Q. An azaacene derivative as promising electron-transport layer for inverted perovskite solar cells. Dyes Pigm. 2021, 192, 109474. [CrossRef]

96. Sasabe, H.; Kido, J. Multifunctional materials in high-performance OLEDs: Challenges for solid-state lighting. Chem. Mater. 2011, 23, 621–630. [CrossRef]

97. Andrulevičiūtė, V.; Leitonas, K.; Volyniuk, D.; Sin, G.; Grazulevičius, J.V.; Getautis, V. TADF versus TTA emission mechanisms in acridan and carbazole-substituted dibenzo[a,c]phenazines: Towards triplet harvesting emitters and hosts. Chem. Eng. J. 2021, 417, 127902. [CrossRef]

98. Yang, W.; Ning, W.; Gong, S.; Yang, C. Deep-red thermally activated delayed fluorescence emitters based on a phenanthroline-containing planar acceptor. Dyes Pigm. 2021, 192, 109474. [CrossRef]

99. Yan, C.; Barlow, S.; Wang, Z.; Yan, H.; Jen, A.K.-Y.; Marder, S.R.; Zhan, X. Non-fullerene acceptors for organic solar cells. Nat. Rev. Mater. 2018, 3, 1–19. [CrossRef]

100. Wadsworth, A.; Moser, M.; Marks, A.; Little, M.S.; Gasparini, N.; Brabec, C.J.; Baran, D.; McCulloch, I. Critical review of the molecular design progress in non-fullerene electron acceptors towards commercially viable organic solar cells. Chem. Soc. Rev. 2019, 48, 1596–1625. [CrossRef]

101. Hahn, S.; Butscher, J.; An, Q.; Jocić, A.; Tverskov, O.; Richter, M.; Feng, X.; Rominger, F.; Vaynzof, Y.; Bunz, U.H.F. Azaarene dimers. Chem. Eur. J. 2019, 25, 7285–7291. [CrossRef]

102. Liang, Z.; Tang, Q.; Mao, R.; Liu, D.; Xu, J.; Miao, Q. The position of nitrogen in N-heteropentacenes matters. J. Mater. Chem. C 2011, 8517, 1–4. [CrossRef]

103. Saliba, M.; Correa-Baena, J.-P.; Grätzel, M.; Hagfeldt, A.; Abate, A. Perovskite solar cells: From the atomic level to film quality and device performance. Angew. Chem. Int. Ed. 2018, 57, 2554–2569. [CrossRef]

104. Boyd, C.C.; Cheacharoen, R.; Leijtens, T.; McGehee, M.D. Understanding degradation mechanisms and improving stability of perovskite photovoltaics. Chem. Rev. 2019, 119, 3418–3451. [CrossRef]

105. Gu, P.-Y.; Wang, N.; Wang, C.; Zhou, Y.; Long, G.; Tian, M.; Chen, W.; Sun, X.W.; Kanatzidis, M.G.; Zhang, Q. Pushing up the efficiency of planar perovskite solar cells to 18.2% with organic small molecules as the electron transport layer. J. Mater. Chem. A 2017, 5, 7339–7344. [CrossRef]

106. Gu, P.-Y.; Wang, N.; Wu, A.; Wang, Z.; Tian, M.; Fu, Z.; Sun, X.W.; Zhang, Q. An azacene derivative as promising electron-transport layer for inverted perovskite solar cells. Chem. Asian J. 2016, 11, 2135–2138. [CrossRef] [PubMed]

107. Wang, N.; Zhao, K.; Ding, T.; Liu, W.; Ahmed, A.S.; Wang, Z.; Tian, M.; Sun, X.W.; Zhang, Q. Improving interfacial charge recombination in planar heterojunction perovskite photovoltaics with small molecule as electron transport layer. Adv. Energy Mater. 2017, 7, 1700522. [CrossRef]

108. Liang, Z.; Tang, Q.; Xu, J.; Miao, Q. Soluble and stable N-heteropentacenes with high field-effect mobility. Adv. Mater. 2011, 23, 1535–1539. [CrossRef] [PubMed]

109. Liu, Y.-Y.; Song, C.-L.; Zeng, W.-J.; Zhou, K.-G.; Shi, Z.-F.; Ma, C.-B.; Yang, F.; Zhang, H.-L.; Gong, X. High and balanced hole and electron mobilities from ambipolar thin-film transistors based on nitrogen-containing oligoacenes. J. Am. Chem. Soc. 2010, 132, 16349–16351. [CrossRef] [PubMed]

110. Song, C.-L.; Ma, C.-B.; Yang, F.; Zeng, W.-J.; Zhang, H.-L.; Gong, X. Synthesis of Tetrachloro-azapentacene as an ambipolar organic semiconductor with high and balanced carrier mobilities. Org. Lett. 2011, 13, 2880–2883. [CrossRef]

111. Liao, H.; Xiao, C.; Ravva, M.K.; Yao, L.; Yu, Y.; Yang, Y.; Zhang, W.; Zhang, L.; Li, Z.; McCulloch, I.; et al. Fused pyrazine-and carbazole-containing azacenes: Synthesis and properties. ChemPlusChem 2019, 84, 1257–1262. [CrossRef]

112. Chu, M.; Fan, J.-X.; Yang, S.; Liu, D.; Ng, C.F.; Dong, H.; Ren, A.-M.; Miao, Q. Halogenated tetrazapentacenes with electron mobility as high as 27.8 cm² V⁻¹ s⁻¹ in solution-processed n-channel organic thin-film transistors. Adv. Mater. 2018, 30, 1803467. [CrossRef]

113. Wu, Y.; Yin, Z.; Xiao, J.; Liu, Y.; Wei, F.; Tan, K.J.; Kloc, C.; Huang, L.; Yan, Q.; Hu, F.; et al. Crystal structure and phototransistor behavior of N-substituted heptacene. ACS Appl. Materials Interfaces 2012, 5, 17–4. [CrossRef]

114. Ganschow, M.; Koser, S.; Hodecker, M.; Rominger, F.; Freudenberg, J.; Dreuw, A.; Bunz, U.H.F. Azaacenes bearing five-membered rings. Chem. Eur. J. 2018, 24, 13667–13675. [CrossRef] [PubMed]

115. He, J.-L.; Kong, F.-C.; Sun, B.; Wang, X.-J.; Tian, Q.-S.; Fan, J.; Liao, L.-S. Highly efficient deep-red TADF organic light-emitting diodes via increasing the acceptor strength of fused polycyclic aromatics. Chem. Eng. J. 2021, 424, 130470. [CrossRef]
115. Hu, Y.; Zhang, Y.; Han, W.; Li, J.; Pu, X.; Wu, D.; Bin, Z.; You, J. Orange–red organic light-emitting diodes with high efficiency and low efficiency roll-off: Boosted by a fused acceptor composed of pyrazine and maleimide. Chem. Eng. J. 2022, 428, 131186. [CrossRef]

116. Maeng, J.H.; Braveenth, R.; Jung, Y.H.; Hwang, S.J.; Lee, H.; Min, H.L.; Kim, J.Y.; Han, C.W.; Kwon, J.H. Efficiency enhancement in orange red thermally activated delayed fluorescence OLEDs by using a rigid di-indolocarbazole donor moiety. Dyes Pigm. 2021, 194, 109980. [CrossRef]

117. Ha, J.M.; Shin, H.B.; Joung, J.F.; Chung, W.J.; Jeong, J.-E.; Kim, S.; Hur, S.H.; Bae, S.-Y.; Kim, J.-Y.; Lee, J.Y.; et al. Rational molecular design of azacene-based narrowband green-emitting fluorophores: Modulation of spectral bandwidth and vibronic transitions. ACS Appl. Mater. Interfaces 2021, 13, 26227–26236. [CrossRef] [PubMed]

118. Ganschow, M.; Koser, S.; Hahn, S.; Rominger, F.; Freudenberg, J.; Bunz, U.H.F. Dibenzobarrelene-based azacenes: Emitters in organic light-emitting diodes. Chem. Eur. J. 2017, 23, 4415–4421. [CrossRef]

119. Li, W.; Pan, Y.; Yao, L.; Liu, H.; Zhang, S.; Wang, C.; Shen, F.; Lu, P.; Yang, B.; Ma, Y. A Hybridized local and charge-transfer excited state for highly efficient OLEDs: Molecular design, spectral character, and full exciton utilization. Adv. Opt. Mater. 2014, 2, 892–901. [CrossRef]

120. Huang, W.; Einzinger, M.; Zhu, T.; Chae, H.S.; Jeon, S.; Ihn, S.-G.; Sim, M.; Kim, S.; Su, M.; Teveroskiy, G.; et al. Molecular design of deep blue thermally activated delayed fluorescence materials employing a homoconjugative triptycene scaffold and dihedral angle tuning. Chem. Mater. 2018, 30, 1462–1466. [CrossRef]

121. Park, Y.; Kim, S.; Lee, J.-H.; Jung, D.H.; Wu, C.-C.; Park, J. New blue-violet emitters based on an indenopyrazine core for OLEDs: Effects of the position of m-terphenyl side group substitution on optical and electroluminescence properties. Org. Electron. 2010, 11, 864–871. [CrossRef]

122. Chen, M.; Perzon, E.; Andersson, M.R.; Marcinkevicius, S.; Jönsson, S.K.M.; Fahlman, M.; Berggren, M. 1 Micron wavelength photo- and electroluminescence from a conjugated polymer. Appl. Phys. Lett. 2004, 84, 3570–3572. [CrossRef]

123. Selzer, F.; Falkenberg, C.; Hamburger, M.; Baumgartner, M.; Müllen, K.; Leo, K.; Riede, M. Improved organic p-i-n type solar cells with n-doped fluorinated hexaazatrinaphthylene derivatives HATNA-F6 and HATNA-F12 as transparent electron transport material. J. Appl. Phys. 2014, 115, 054515. [CrossRef]

124. Zhao, D.; Zhu, Z.; Kuo, M.-Y.; Chueh, C.-C.; Jen, A.K.-Y. Hexaazatrinaphthylene derivatives: Efficient Electron-transporting materials with tunable energy levels for inverted perovskite solar cells. Angew. Chem. Int. Ed. 2016, 55, 8999–9003. [CrossRef] [PubMed]

125. Zhu, Z.; Zhao, D.; Chueh, C.-C.; Shi, X.; Li, Z.; Jen, A.K.-Y. Highly efficient and stable perovskite solar cells enabled by all-crosslinked charge-transporting networks. Joule 2018, 2, 168–183. [CrossRef]

126. Lindner, B.D.; Paulus, F.; Appleton, A.L.; Schaffroth, M.; Engelhart, J.U.; Schelkle, K.M.; Tverskoy, G.; Rominger, F.; Hamburger, M.; Bunz, U.H.F. Electron-transporting phenazinothiadiazoles with engineered microstructure. J. Mater. Chem. C 2014, 2, 9609–9612. [CrossRef]

127. Thiadiazolo-Azaacenes-Müller-2019-Chemistry–A European Journal-Wiley Online Library. Available online: https://chemistry-europe.onlinelibrary.wiley.com/doi/full/10.1002/chem.201900462 (accessed on 20 August 2021).

128. Xu, X.; Yao, Y.; Shan, B.; Gu, X.; Liu, D.; Liu, J.; Xu, J.; Zhao, N.; Hu, W.; Miao, Q. Electron mobility exceeding 10 cm2 V−1 s−1 and band-like charge transport in solution-processed n-channel organic thin-film transistors. Adv. Mater. 2016, 28, 5276–5283. [CrossRef] [PubMed]

129. Paulus, F.; Forz, M.; Schaffroth, M.; Rominger, F.; Leineweber, A.; Vaynzofo, Y.; Bunz, U.H.F. Side-group engineering: The influence of norbornadienyl substituents on the properties of ethynylated pentacene and tetrazapentacene. Org. Electron. 2016, 33, 102–109. [CrossRef]

130. Reiss, H.; Ji, L.; Han, J.; Koser, S.; Tverskoy, O.; Freudenberg, J.; Hinkel, F.; Moos, M.; Friedrich, A.; Krummenacher, I.; et al. Bromination improves the electron mobility of tetrazapentacene. Angew. Chem. Int. Ed. 2018, 57, 9543–9547. [CrossRef] [PubMed]

131. Brosius, V.; Weigold, S.; Hippchen, N.; Rominger, F.; Freudenberg, J.; Bunz, U.H.F. Diindenopyrazines: Electron-deficient arenes. Chem. Eur. J. 2021, 27, 10001–10005. [CrossRef]

132. Gu, P.-Y.; Wang, Z.; Liu, G.; Yao, H.; Wang, Z.; Li, Y.; Zhu, J.; Li, S.; Zhang, Q. Synthesis, full characterization, and field effect transistor behavior of a stable pyrene-fused N-heteroacene with twelve linearly annulated six-membered rings. Chem. Mater. 2017, 29, 4172–4175. [CrossRef]

133. Endres, A.H.; Schaffroth, M.; Paulus, F.; Reiss, H.; Wadepohl, H.; Rominger, F.; Krämer, R.; Bunz, U.H.F. Coronene-containing N-heteroarenes: 13 rings in a row. J. Am. Chem. Soc. 2016, 138, 1792–1795. [CrossRef]

134. Rohnacher, V.; Benneckendorf, F.S.; Münch, M.; Sauter, E.; Asyuda, A.; Barf, M.-M.; Tisserant, J.-N.; Hillebrandt, S.; Rominger, F.; Jänsch, D.; et al. Functionalized tetrapodal diazatriptycenes for electrostatic dipole engineering in N-type organic thin film transistors. Adv. Mater. Technol. 2021, 6, 2000300. [CrossRef]

135. Gu, P.-Y.; Wang, Z.; Xiao, F.-X.; Lin, Z.; Song, R.; Xu, Q.-F.; Lu, J.-M.; Liu, B.; Zhang, Q. An ambipolar azacene as a stable photocathode for metal-free light-driven water reduction. Mater. Chem. Front. 2017, 1, 495–498. [CrossRef]

136. Wang, C.; Wang, J.; Li, P.; Gao, J.; Tan, Y.; Xiong, W.; Hu, B.; Lee, S.; Zhao, Y.; Zhang, Q. Synthesis, characterization, and non-volatile memory device application of an N-substituted heteroacene. Chem. Asian J. 2014, 9, 779–783. [CrossRef] [PubMed]
137. Zhao, K.; Yu, F.; Liu, W.; Huang, Y.; Said, A.A.; Li, Y.; Zhang, Q. Unexpected synthesis, properties, and nonvolatile memory device application of imidazole-fused azacenes. *J. Org. Chem.* **2020**, *85*, 101–107. [CrossRef] [PubMed]

138. Li, G.; Zheng, K.; Wang, C.; Leck, K.S.; Hu, F.; Sun, X.W.; Zhang, Q. Synthesis and nonvolatile memory behaviors of dioxate-traazapentacene derivatives. *ACS Appl. Mater. Interfaces* **2013**, *5*, 6458–6462. [CrossRef] [PubMed]

139. De Girolamo, J.; Reiss, P.; Pron, A. Hybrid materials from diaminopyrimidine-functionalized poly(hexylthiophene) and thymine-capped CdSe nanocrystals: Part II—Hydrogen bond assisted layer-by-layer molecular level processing. *J. Phys. Chem. C* **2008**, *112*, 8797–8801. [CrossRef]