Molecules 2000, 5, 1331-1371



http://www.mdpi.org

Organic Iodine(I, III, and V) Chemistry: 10 Years of Development at the Medical University of Warsaw, Poland[†]

Lech Skulski*

Chair and Laboratory of Organic Chemistry, Faculty of Pharmacy, Medical University of Warsaw, 1 Banacha Street, PL 02-097 Warsaw, Poland. Tel./fax: +(4822) 8226843.

[†] Presented as Review A0004 at the 4th Electronic Conference on Synthetic Organic Chemistry (ECSOC-4), September 1-30, 2000.

*To whom correspondence should be addressed. E-mail: lskulski@farm.amwaw.edu.pl

Received: 16 November 2000 / Accepted: 16 November 2000 / Published: 20 December 2000

Abstract: This review reports some *novel* (or considerably improved) methods for the synthesis of aromatic iodides, (dichloroiodo)arenes, (diacetoxyiodo)arenes, iodylarenes and diaryliodonium salts, as well as some facile, *oxidative* anion metatheses in *crude* diaryliodonium halides and, for comparison, potassium halides. All these new results were obtained in our laboratory over the past decade (1990-2000). A full list of our papers dealing with the organic iodine(I, III and V) chemistry, covering exlusively the *aromatic* derivatives, is also provided.

Keywords: aromatic oxidative iodination, iodoarenes, aromatic hypervalent iodine compounds, (dichloroiodo)arenes, (diacetoxyiodo)arenes, iodylarenes, diaryliodonium salts, oxidative anion metatheses in ionic halides

Contents

- 1. Introduction
- 2. Syntheses of Aromatic Iodides
- 3. Syntheses of (Dichloroiodo)arenes
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- 4. Syntheses od (Diacetoxyiodo)arenes
- 5. Syntheses of Iodylarenes
- 6. Syntheses of Diaryliodonium Salts
- 7. Anion Metatheses in Diaryliodonium Halides
- 8. Conclusions

1. Introduction

In the years 1980-1990 our research group had been mainly interested in the chemistry of aromatic, symmetric and unsymmetric organomercurials; see Refs. 22-33. In particular, we successfully synthesized a number of novel organomercurials derived from lactamic heterocycles as well as we discovered several novel cyano- and halo-demercuration reactions; this resulted in many effective syntheses of cyano or halo derivatives of those aromatics, whose corresponding organomercurials had been used in the said novel demercuration reactions. However, the quickly growing number of literature reports and reviews on the unique synthetic possibilities offered by organic hypervalent iodine reagents/compounds had been so promising and alluring that in 1990 we definitely decided that nearly all our future research work should be directed towards this area. Since all our previous research activity had been concerned with aromatic compounds, all our subsequent studies have also been limited to aromatic iodine(III or V) compounds. Additionally, we put special emphasis on the development of novel, easy and effective oxidative iodination procedures, applicable for numerous aromatics (both activated and deactivated), since the resulting iodo derivatives would then applied by us in our subsequent syntheses of the corresponding aromatic hypervalent iodine compounds. So far, mainly (dichloroiodo)arenes, (diacetoxyiodo)arenes, iodylarenes, and aromatic iodonium salts have been synthesized and studied by us, using novel (or considerably improved) methods. This short review presents mainly our own contributions to the chemistry of aromatic iodine(I, III and V) compounds over the past decade (1990-2000). At the end of this review a full list of our papers and Ph.D. theses dealing with this topic is provided (see Refs. 1-18).

2. Syntheses of Aromatic Iodides

Aromatic iodides, known for ca.150 years, are generally more reactive than the respective bromides and chlorides. There is a considerable number of different methods, direct or indirect, for their synthesis, and they are widely used in chemical laboratories and, sometimes, also in chemical industry and medicine [19]. Moreover, they are able to form a large variety of stable, aromatic polyvalent iodine compounds, which have found increasing applications in modern organic synthesis [21]. 2.1. Our earlier results: aromatic iodides from aromatic organomercurials [22-33]

It is known [20] that aromatic iodides may be obtained *indirectly* from both symmetric and unsymmetric aromatic organomercurials, by reacting them with either hot aq. KI₃ solutions or with hot solutions of diiodine in acetonitrile, DMF, etc. We had repeatedly applied the "classic" *iododemercuration* reactions to better elucidate the chemical structures of organomercurials prepared by us in various ways. Thus, we synthesized (the yields given in brackets) the following *aromatic iodides*:

i) 8-iodocaffeine (95%) from both 8-(acetoxymercurio)caffeine and 8,8'-mercuriobis(caffeine) [22];

ii) *5-iodouracil* (60 or 93%, resp.) from either 1-acetyl-5-(trifluoroacetoxymercurio)uracil or 5,5'-mercuriobis(uracil) [23];

iii) 8-iodotheophylline (95%) from 8-(trifluoroacetoxymercurio)theophylline, and 8-iodotheobromine (95%) from 8-(trifluoroacetoxymercurio)theobromine [24];

iv) 2-iodofuran (60%) from 2,2'-difurylmercury, and 2-iodothiophene (65%) from 2,2'-dithienylmercury [25];

v) *5-iodo-1,3-dimethyluracil* (87.5 or 82.5%, resp.) from either 5-(acetoxymercurio)-1,3-dimethyluracil or 5,5'-mercuriobis(1,3-dimethyluracil) [27];

vi) *5-iodo-2,4-dimethoxypyrimidine* (58 or 52%, resp.) from either 5-(acetoxymercurio)-2,4-dimethoxypyrimidine or 5,5'-mercuriobis(2,4-dimethoxypyrimidine); the acid hydrolysis of pure 5-iodo-2,4-dimethoxypyrimidine afforded *5-iodouracil* (69%) [28];

vii) *6-iodo-2,3-diphenyl-5-methoxybenzofuran* (70%) from both 6-(acetoxymercurio)-2,3-diphenyl-5-methoxybenzofuran and 6,6'-mercuriobis(2,3-diphenyl-5-methoxybenzofuran) [29];

viii) *iodobenzene* (59%) from (chloromercurio)benzene, and *8-iodocaffeine* (65%) from 8-(chloromercurio)caffeine [30];

ix) *8-iodotheobromine* (97%) from 1,8-bis(acetoxydimercurio)theobromine; the latter compound was the first *stable* organic derivative of **mercury(I)** [32];

x) *5-iodouracil* (61% crude yield) from 5-(chloromercurio)uracil, and *2-ethylthio-4-hydroksy-5-iodopyrimidine* (64% crude yield) from 2-ethylthio-4-hydroxypyrimidine raw mercurial [33].

We also applied twice our *novel* iodo-demercuration method, by reacting two *symmetric* organomercurials with hot ethanolic solutions of cyanogen iodide, ICN. In this way we synthesized:

xi) 8-iodocaffeine (90%) from 8,8'-mercuriobis(caffeine) [22], and

xii) iodobenzene (72%) from diphenylmercury [26].

2.2. Aromatic Iodides from Some Highly Activated Aromatics with Lead(IV) Acetate as Oxidant [2]

In a very brief communication, without an experimental part, Soviet authors [34] have reported that on treating PhR (R = H, F, Cl, Br, I, Me, CF₃) with diiodine and Pb(OAc)₄ dissolved in *costly* CF₃CO₂H, the corresponding iodides are formed in 85-90% yields, as determined by GLC. Stoichiometric ratios of Pb⁴⁺/I₂/Ar-H = 1:1:2 were mostly applied. They assumed that both *iodine(I)*

1334

and *iodine(III)* species would be the iodinating agents. A possible formation of intermediate $ArI(OCOCF_3)_2$ from Ar-H was claimed based only on the evidence of the reaction of PhCl (no yield given) with a Pb⁴⁺/I₂ (3:1) system, in which the following stoichiometry was obeyed:

 $I_2 + 3Pb^{4+} \rightarrow 2I^{3+} + 3Pb^{2+}$

However, details of their iodination method have never been published.

In our first work [2] we carried out the oxidative aromatic iodination of eleven *highly activated* arenes and heterocycles in the presence of excess (10%) pure Pb(OAc)₄ in glacial acetic acid, used as the *inexpensive* solvent of choice:

 $Pb(OAc)_4 + I_2 \rightarrow Pb(OAc)_2 + 2AcOI (transient acetyl hypoiodite)$ 2Ar-H + 2AcOI \rightarrow 2ArI + 2AcOH, hence:

$$2 \operatorname{Ar-H} + I_2 + \operatorname{Pb}(\operatorname{OAc})_4 \xrightarrow{\text{r.t., } 0.5 - 12 \text{ h}} 2 \operatorname{Ar-I} + \operatorname{Pb}(\operatorname{OAc})_2 + 2 \operatorname{AcOH}$$
(Eq. 1).

It is of interest that when using hot $Pb(OAc)_4$ solutions prepared *in situ* from finely powdered, commercial minimum, Pb_3O_4 , prior to the iodination, i.e.:

$$Pb_{3}O_{4} + 4 Ac_{2}O \xrightarrow{55-60 \text{ oC}, 15-30 \text{ min}}_{\text{nearly quantitatively}} \rightarrow Pb(OAc)_{4} + 2 Pb(OAc)_{2}$$

then the iodination yields were often somewhat better than those obtained with the pure oxidant. Also 4-bromo- and 2,4-dibromoanisoles were obtained from anisole by the respective *oxidative brominations*. *We failed* to iodinate benzene, toluene, 1,3-xylene and several less active arenes or heteroaromatics; for more information see Ref. 2.

2.3. Aromatic Iodides from Activated or Deactivated Arenes with Chromium(VI) Oxide as Oxidant [8]

Japanese chemists [35] iodinated benzene with diiodine, in pressurized air and oxygen gas and in acidic solvents using twenty four metal salts or oxides as *catalysts*; they mentioned also *a chromate* and MnO_2 (see Section 2.5). The iodination rate was dependent on the acidity of the solvent applied: $CF_3SO_3H > CH_3SO_3H > CF_3COOH >> CH_3COOH$. PhI was obtained in excellent yield (91.9%) when NaVO₃ and CF_3SO_3H were used. According to them, the combination of strong acidic solvent, oxygen gas, and oxidizing catalyst are indispensable in the iodination reaction, whereas the kind of metal counter ion used does not seem to be important. Later, Soviet chemists [36] used *chromium (VI)* salts

for the oxidative aromatic iodination of benzene and toluene with diiodine in a mixture of CCl₄ with acetic acid containing H₂SO₄, using four dichromates as the oxidants. They qualitatively arranged the oxidizing activity of the dichromates as follows: $K_2Cr_2O_7 < Na_2Cr_2O_7 < Li_2Cr_2O_7 < (NH_4)_2Cr_2O_7$. We assume that their failure to iodinate PhCOOH and PhNO₂ was mainly due **to the presence of WATER** in the reaction mixtures: they always used *water* acidified with H₂SO₄ for prior dissolution of the dichromates. It is known [37] that *water*, due to its high hydration power, diminishes the oxidizing activity of inorganic (and organic) oxidants. In contrast, this activity is increased when *the overall acidity* of the oxidizing mixtures rises to some optimum level. When this level is considerably exceeded, then the iodination yields may drop strikingly, due to a predominant *protonation* of the reacted aromatic molecules.

Note. Starting from our second paper in the series [8], in all our subsequent studies dealing with the oxidative aromatic iodination of various arenes/heteroarenes, the above general considerations have always been taken into account. The favorable *anhydrous* conditions were simply attained using mixtures of glacial acetic acid with *acetic anhydride* as the cheap solvents of choice. The optimum overall acidities of the iodinating mixtures, which are *different* for each arene/heteroarene iodinated, were attained by dropwise addition to the cooled reaction mixtures of *widely varied* quantities of conc. (98%) sulfuric acid, established experimentally *in each case*; see Experimental parts in our papers [8, 11, 15, 17, 18c,d] for more details.

For benzene, halobenzenes, and several *activated* arenes, the following reaction mechanism and general reaction stoichiometry were assumed and applied:

$$3I_{2} + 2Cr(VI) \rightarrow 6I^{+} + 2Cr(III) (a \text{ preliminary stoichiometry})$$

$$6Ar-H + 6I^{+} \rightarrow 6ArI + 6H^{+}, \text{ hence:}$$

$$6 \text{ Ar-H} + 3 I_{2} + 2 \text{ CrO3} + 6 \text{ Ac}_{2}O + 3 \text{ H}_{2}\text{SO4} \longrightarrow$$

$$AcOH/Ac_{2}O$$

$$55-65 \text{ oC}, 0.5 - 1.5 \text{ h} \qquad 6 \text{ Ar-I} + Cr_{2}(SO_{4})_{3} + 12 \text{ AcOH} \qquad (Eq. 2).$$

In spite of many attempts, working under widely varied reaction conditions, in practice we were *unable* to iodinate nitrobenzene. For a less strongly deactivated arene, benzoic acid, the final yield was low, i.e. 30%. Hence, for the first time in our research work, we *deliberately* changed the proportions of the reactants and, consequently, the *reaction mechanism*, by carrying out the oxidative aromatic iodination reactions as follows [the same general approach has later been used in all our subsequent iodination procedures applicable for *deactivated* aromatics, *vide infra*]:

$$I_{2} + 2Cr(VI) \rightarrow 2I^{3+} + 2Cr(III) (a \ preliminary \ stoichiometry)$$

$$\underbrace{2Ar-H + 2I^{3+} + 2H_{2}SO_{4} \rightarrow 2ArISO_{4} + 6H^{+}, \ hence:}{2 \ Ar-H + I_{2} + 2 \ CrO_{3} + 6 \ Ac_{2}O + 5 \ H_{2}SO_{4} \longrightarrow}$$

$$AcOH/Ac_{2}O$$

$$\underbrace{r.t.- 60 \ ^{o}C, \ 4-7 \ h}{2 \ ArISO_{4} (not \ isolated) + Cr_{2}(SO_{4})_{3} + 12 \ AcOH} \qquad (Eq. 3).$$

After completing the above reactions, the resulting (final) reaction mixtures containing the assumed organic *iodine(III)* intermediates, ArISO₄, were poured into excess aq. Na_2SO_3 solutions to give the corresponding crude iodoarenes, ArI, as follows:

$$2ArISO_4 (not isolated) + 2Na_2SO_3 + 2H_2O \rightarrow 2ArI + 4NaHSO_4$$
(Eq. 4)

After the isolation and purification of crude ArI, we obtained pure 3-IC₆H₄COOH (86%), 3-IC₆H₄COOMe (66%), 3-IC₆H₄COOEt (64%), 3-IC₆H₄CONH₂ (88%), 3-IC₆H₄SO₂NH₂ (74%), and 3-IC₆H₄NO₂ (71%). Similarly, it was possible to obtain some pure diiodinated products, viz. 3,3'-diiodobenzophenone (57%), 3,3'-diiododibenzoyl (72%), 3,3'-diiododiphenyl sulfone (83%), and 3,5-diiodobenzoic acid (31%). These high iodination yields we achieved with *deactivated* arenes were due to the fact that the respective arenes had been reacted with some preponderant, *more electrophilic* transient *iodine(III)* species, I^{3+} , formed momentarily in the reaction mixtures prepared in agreement with Eq. 3. Contrariwise, for benzene, halobenzenes, and *activated* arenes, the best iodination yields were attained with the reaction mixtures prepared in agreement with Eq. 2, where some *less electrophilic*, transient *iodine(I)* species, I^{+} , had evidently been predominant.

When *benzene* was iodinated according to Eq. 3, the yield of PhI was low (10%). However, from the collected aqueous layers we precipitated out a yellow solid with excess aq. KBr. After recrystallization from MeOH is was identified as *diphenyliodonium bromide* (44%). We commented on this fact as follows [8]: "This novel "one-pot" method of preparing the symmetric diaryliodonium salts is presently being improved and extended [70], and will be published soon". However, publication of this method has been delayed.

A more detailed discussion on the transient I^+ or I^{3+} species that probably exist in the iodinating mixtures prepared in agreement either with Eq. 2 or with Eq. 3, was subsequently submitted in our last paper of the series [17]; see Section 2.6. The *varied quantities* of conc. H₂SO₄ (established experimentally *for each* of the arenes iodinated) added dropwise to the cooled reaction mixtures clearly depended on relative reactivities (and basicities) of the arenes investigated. The more deactivated an arene was, more conc. H₂SO₄ had to be added to catalyze better the iodination, and to possibly increase the concentration of *still more reactive* either *iodine(II)* or *iodine(III)* transient species:

$$I_2(SO_4) \xrightarrow{+ H_2SO_4} 2IOSO_3H \xrightarrow{- } 2I^{\oplus} + 2 HSO4^{\ominus}$$

(all denoted above as \mathbf{I}^+)

$$I_{2}(SO_{4})_{3} \xrightarrow{+ H_{2}SO_{4}} 2 I(SO_{4})OSO_{3}H \xrightarrow{- 2} 2 I^{\oplus}(SO_{4}) + 2 HSO_{4}^{\ominus}$$
$$\xrightarrow{+ H_{2}SO_{4}} 2 I^{\oplus}(OSO_{3}H)_{2} + 2 HSO_{4}^{\ominus}$$

(all denoted above as I^{3+})

In the strongly acidic reaction mixtures discussed, some fairly stable organic *iodine(III)* intermediates, $ArISO_4$ (see Eq. 3), are, probably *quickly equilibrating* with some other intermediates, viz. $ArI(OSO_3H)_2$ (which, in fact, were solely shown in Ref. [8]) on adding more H₂SO₄. For the sake of simplicity, *only one kind of them* was shown and discussed in all our papers. Their real existence in the reaction mixtures has been experimentally demonstrated in our newest work [18c]; see Section 4.1.

2.4. Aromatic Iodides from Deactivated Arenes with Potassium Permanganate as Oxidant [11]

Chaikovski and Novikov [38] oxidatively iodinated benzene and various activated and deactivated derivatives, and some aromatic hydrocarbons, using stoichiometric quantities of $KMnO_4$ in strongly acidic media. To explain the results, they proposed the following assumed reaction mechanism:

$$\begin{split} & 5I_2 + 2KMnO_4 + 3H_2SO_4 + 2H_2O = K_2SO_4 + 2MnSO_4 + 10HIO \\ & HIO + H_2SO_4 \rightarrow H_2^+OI + HSO_4^- \\ & H_2^+IO \rightarrow H_2O + \mathbf{I}^+ \end{split}$$

This shows, in their opinion, that a protonated form of hypoiodic acid, H_2^+OI , generated from diiodine by the action of KMnO₄ in strongly acidic solutions, is the main source of active, transient *iodine(I)* species, **I**⁺, which then react momentarily with the iodinated aromatics.

They worked out two simple procedures, carrying out the iodination reactions (at 100-115°C, for 1-5 h) using the following systems: *Procedure 1*: $I_2/KMnO_4/glacial AcOH/conc. H_2SO_4$ or *Procedure 2*: $I_2/KMnO_4/80\%$ AcOH/conc. H_2SO_4. Generally, the corresponding mono- or diiodinated products were obtained in 26-99% yields - hence, we made no attempt to improve their *Procedure 2*, suitable for aromatic hydrocarbons and *activated* arenes. But in the *Procedure 1* they used, quite purposefully, *an industrial grade glacial acetic acid* - probably containing a considerable admixture of water which, due to its high hydration power, diminishes the oxidizing activity of inorganic oxidants [37]. Hence, on iodinating PhCOOH and 9,10-phenanthrenequinone they obtained the iodinated products in only moderate yields: $3\text{-IC}_6\text{H}_4\text{COOH}$ (49%) and 2,7-diiodo-9,10-phenanthrenequinone (35%). Taking into account our former experiments [8] explained in Section 2.3, we were able *to improve considerably* said *Procedure 1*, suitable for the iodination of *deactivated* arenes, by carrying out the iodination reactions under *anhydrous* conditions, and by applying the following reaction stoichiometry, which strongly favors the formation of more electrophilic, transient *iodine(III)* species, viz.

 $5I_2 + 6Mn(VII) \rightarrow 10I^{3+} + 6Mn(II)$ (a preliminary stoichiometry)

$$\underbrace{\frac{10\text{Ar-H} + 10\mathbf{I}^{3+} + 10\text{H}_2\text{SO}_4 \rightarrow 10\text{ArISO}_4 + 20\text{H}^+, \text{ hence:}}{2 \text{ Ar-H} + I_2 + 1.2 \text{ KMnO4} + 4.8 \text{ Ac}_2\text{O} + 3.8 \text{ H}_2\text{SO4}} \longrightarrow$$

$$\underbrace{\text{AcOH/Ac}_2\text{O}}{35 \text{ °C}, 1 \text{ h}} 2 \text{ ArISO}_4 (not isolated) + 1.2 \text{ MnSO}_4 + 0.6 \text{ K}_2\text{SO}_4 + 9.6 \text{ AcOH}$$
(Eq. 5).

After pouring the final reaction mixtures containing fairly stable organic ArISO₄ intermediates, into excess aq. Na₂SO₃ solutions (see Eq. 4 in Section 2.3), and the separation and purification of the crude products, we thus obtained 3-IC₆H₄COOH (78%), 3-IC₆H₄COOMe (83%) and 3-IC₆H₄COOEt (73%) as well as 2,7-diiodo-9,10-phenanthrenequinone (85%), by lowering the reaction temperature *to* 35°C, and shortening the reaction time *to one hour*.

It is necessary to add that Chinese chemists from Taiwan [39] iodinated various arylamines with a homogeneous mixture of hydroiodic acid and **KMnO**₄ in acetonitrile, obtaining *para*-substituted products in high yields (71-78%) within twelve hours at room temperature. This reagent is highly selective for arylamines, but unsuitable for phenols, which undergo oxidative coupling easily in the presence of electron transfer agents.

2.5. Aromatic Iodides from Activated and Deactivated Arenes with Various Brands of Manganese(IV) Oxide as Oxidants [11, 18d]

Activated MnO_2 is a common and easily handled oxidant, and its many applications in organic synthesis have been known for a long time [40]. Moreover, Mn(II) salts remaining in the residues after the oxidative iodination reactions are *less toxic* as compared with Pb(II) or Cr(III) salts, the byproducts in our former iodination experiments [2, 8]. Starting our iodination experiments with **MnO₂**, we checked out *various brands* of this oxidant experimentally. Ordinary commercial MnO₂ was not applicable. The *best* iodination yields were obtained with the *activated* MnO₂ freshly prepared prior to use [41]. An *activated* commercial product (Aldrich AMD, suitable for organic reactions, ca. 85% MnO₂) gave lower iodination yields by ca. 5-10%. The so-called Chemical Manganese Dioxide (CMD, suitable for use in batteries) has not been studied by us until recently [18d]; our newest results are briefly discussed below, at the end of this Section. For benzene, toluene, and four halobenzenes, the iodination reactions were carried out according to the following stoichiometry, which favors the preponderant formation of transient iodine(I) species, viz.

$$I_2 + Mn(IV) \rightarrow 2I^+ + Mn(II)$$
 (a preliminary stoichiometry)

$$\frac{2\text{Ar-H} + 2\mathbf{I}^{+} \rightarrow 2\text{ArI} + 2\text{H}^{+}, \text{ hence:}}{2 \text{ Ar-H} + I_{2} + \text{MnO}_{2} + 2 \text{ Ac}_{2}\text{O} + \text{H}_{2}\text{SO}_{4} \longrightarrow}$$

$$\frac{\text{AcOH/Ac}_{2}\text{O}}{\frac{\text{r.t., 2 h; 45-55 °C, 2 h}}{62-75\%} 2 \text{ Ar-I} + \text{MnSO}_{4} + 4 \text{ AcOH}} \qquad (Eq. 6).$$

Similarly, we obtained 1,4-diiodobenzene (66%) from benzene, when we halved the amount of Ar-H (i.e. benzene) in Eq. 6. For the oxidative iodination of *deactivated* arenes, the deliberately changed general reaction stoichiometry was applied, which favors the preponderant formation of transient *iodine(III)* species, viz.

$$I_{2} + 3Mn(IV) \rightarrow 2I^{3+} + 3Mn(II) (a \text{ preliminary stoichiometry})$$

$$\underbrace{2Ar-H + 2I^{3+} + 2H_{2}SO_{4} \rightarrow 2ArISO_{4} + 4H^{+}, \text{ hence:}}_{AcOH/Ac_{2}O} + 6 Ac_{2}O + 5 H_{2}SO_{4} \longrightarrow \\AcOH/Ac_{2}O \xrightarrow{\text{r.t., 2 h; 45-55 °C, 2-9 h}} 2 ArISO_{4} (not isolated) + 3 MnSO_{4} + 12 AcOH (Eq. 7).$$

After pouring the final reaction mixtures, containing the fairly stable organic ArISO₄ intermediates, into excess aq. Na₂SO₃ solutions (see Eq. 4 in section 2.3), and purification of the crude products, we obtained: $3-IC_6H_4COOH$ (87%), $3-IC_6H_4COOMe$ (89%), $3-IC_6H_4COOEt$ (79%), $3-IC_6H_4CONH_2$ (63%), and $3-IC_6H_4NO_2$ (73%). As above, we also obtained the diiodinated products: 3,3'-diiodobenzophenone (78%) and 2,7-diiodo-9,10-phenanthrenequinone (72%).

When *benzene* was iodinated according to Eq. 7, the yield of PhI was negligible, but from the collected aqueous layers we precipitated a yellow solid with excess aq. KI. After its recrystallization from MeOH, it was identified as *diphenyliodonium iodide* (43%).

Japanese chemists [42] have used as an oxidant the so-called Chemical Manganese Dioxide (CMD, Wako product, min. 75% MnO₂; its price in Poland: 378.00 DM/500 g + freight cost 60.00 DM) in numerous selective oxidations of various classes of organic compounds; these reactions often proceeded in nearly quantitative yields under relatively mild conditions. In some cases, the Wako product proved to be much superior to usual active MnO₂ (AMD) commercially available from

Aldrich, Fluka, Merck, Nakarei and Wako companies. Hence, we decided to check its application in the oxidative aromatic iodination reactions presented in Eqs. 6 and 7, respectively. With the Aldrich CMD (90+% MnO₂; its price in Poland: 32.30 DM/500 g), the final yields for the purified ArI were practically the same as those obtained with the Aldrich AMD (its price in Poland: 107.30 DM/500 g). In contrast, with the said Wako CMD the crude iodinated products, isolated by more laborious methods, were dark-colored and heavily contaminated, which resulted *in considerable losses* during their troublesome purification. Though the *self-prepared* AMD (*vide supra*) is, in fact, somewhat more efficient, its tedious preparation is time-consuming and relatively expensive, so we would recommend the application of the Aldrich CMD as the oxidant of choice in oxidative aromatic iodination reactions, since is has satisfactory activity and is *notably less costly*.

2.6. Aromatic Iodides from Activated and Deactivated Arenes with Sodium Periodate or Sodium Iodate as Oxidants [17]

So far, a number of authors have used periodic acid or its salts, and iodic acid or its salts to oxidatively iodinate a variety of *activated* aromatics [19b, 43], including *halobenzenes* [44]. Only the most costly I₂O₅ has scarcely ever been used until recently [45, 46]. Suzuki [47] concluded his extensive studies by stating that the oxidative iodination with periodic acid is "the most convenient method for preparation of mono- or diiodo derivatives from various polyalkyl-benzenes in high yields". Merkushev [19b] summarized the literature results as follows: "The iodination in the presence of iodic acid and periodic acid is accelerated considerably by the addition of *sulfuric acid* and, in some cases, of **water**. These *mild oxidants* are widely used in the iodination reactions of polynuclear aromatic compounds".

Periodic acid, NaIO₄, iodic acid, and NaIO₃ are commercially available, with the two sodium salts being *notably cheaper* [48]. These salts in solutions or suspensions *acidified* with excess conc. H_2SO_4 would momentarily form *in situ* either HIO₄ or HIO₃, respectively. Hence, they were used as such in all our oxidative iodination experiments [17] discussed below.

For benzene, toluene, four halobenzenes, 4-nitroanisole, and N,N-dimethylaniline their oxidative iodination reactions were carried out by us according to the following stoichiometry, which strongly favors the preponderant formation of less electrophilic, transient *iodine(I)* species, viz.

$$\begin{array}{l} 3I_2 + I(VII) \rightarrow 7I^+ (a \ preliminary \ stoichiometry) \\ \hline \underline{7Ar-H + 7I^+ \rightarrow 7ArI + 7H^+, \ hence:} \\ 14 \ Ar-H + 6 \ I_2 + 2 \ \textbf{NaIO4} + 8 \ Ac_2O + H_2SO_4 \longrightarrow \\ \hline AcOH/Ac_2O \\ \hline \underline{mostly \ r.t.; \ 4-5 \ h} \\ \hline 55-76\% \end{array} 14 \ \textbf{Ar-I} + Na_2SO_4 + 16 \ AcOH \end{array}$$
(Eq. 8).

The least reactive $4-O_2NC_6H_4OMe$ was at first stirred at room temperature for one hour, but its iodination was completed by a further stirring for one hour at 45°C. The most reactive PhNMe₂ was iodinated in 65% yield by stirring at +5°C for only one hour; for some reasons, *we failed* to iodinate likewise PhNH₂ and PhNEt₂. When we halved the amount of Ar-H (i.e. benzene) added to the reaction mixture shown in Eq. 8, then we obtained the purified 1,4-C₆H₄I₂ in 85% yield.

For nitrobenzene, 4-nitroanisole, (trifluoromethyl)benzene, benzoic acid and its methyl and ethyl esters, 4-toluic acid, 4-chlorobenzaldehyde [however, *we failed* to iodinate likewise benzaldehyde, benzonitrile, and benzamide] as well as (for comparison) iodobenzene, bromobenzene, and chlorobenzene, we deliberately applied the following general reaction stoichiometry, which strongly favors the preponderant formation of more electrophilic, transient *iodine(III)* species, viz.

 $2I_{2} + 3I(VII) \rightarrow 7I^{3+} (a \text{ preliminary stoichiometry})$ $\underline{7Ar-H + 7I^{3+} + 7H_{2}SO_{4} \rightarrow 7ArISO_{4} + 21H^{+}, \text{ hence:}}$ $14 \text{ Ar-H} + 4 I_{2} + 6 \text{ NaIO}_{4} + 24 \text{ Ac}_{2}O + 17 \text{ H}_{2}SO_{4} \longrightarrow$ $AcOH/Ac_{2}O$ $\underline{mostly r.t.; 3-8 h} = 14 \text{ ArISO}_{4} (not \text{ isolated}) + 3 \text{ Na}_{2}SO_{4} + 48 \text{ AcOH} \qquad (Eq. 9).$

After completing the above reactions, the resulting (final) reaction mixtures, containing the fairly stable organic *iodine(III)* intermediates, ArISO₄, were poured into excess aq. Na₂SO₃ solutions to give the corresponding crude iodoarenes, ArI (see Eq. 4 in Section 2.3). After the purification of crude ArI, we obtained the eleven purified monoiodinated products in good or excellent yields (56-95%). However, when *benzene* was likewise monoiodinated, rather impure PhI was obtained in only 43% yield. This novel procedure is also suitable for the oxidative diiodination of *deactivated* arenes: the purified 3,3'-diiodobenzophenone was obtained in 51% yield.

Finally, using the appropriate $I_2/NaIO_3/Ac_2O/AcOH/conc$. H_2SO_4 anhydrous system - equally suitable for the oxidative iodination of *deactivated* arenes - we monoiodinated nitrobenzene, benzoic acid, and 4-toluic acid, as well as (for comparison) bromobenzene, according to the general reaction stoichiometry, which strongly favors the preponderant formation of more electrophilic, transient *iodine(III)* species, viz.

 $I_{2} + 3I(V) \rightarrow 5I^{3+} (a \text{ preliminary stoichiometry})$ $5Ar-H + 5I^{3+} + 5H_{2}SO_{4} \rightarrow 5ArISO_{4} + 15H^{+}, \text{ hence:}$ $10 \text{ Ar-H} + 2 I_{2} + 6 \text{ NaIO3} + 18 \text{ Ac}_{2}O + 13 \text{ H}_{2}SO_{4} \longrightarrow$ $AcOH/Ac_{2}O$ $mostly \text{ r.t.; } 3-7 \text{ h} 10 \text{ ArISO4} (not isolated) + 3 \text{ Na}_{2}SO_{4} + 36 \text{ AcOH}$ (Eq. 10).

After completing the main iodination reactions within 3-5 h at room temperature (only PhNO₂ was stirred at r.t. for 4 h, and then at 65° C for 3 h), the strongly acidic reaction mixtures were poured into excess aq. Na₂SO₃ solutions (see Eq. 4 in Section 2.3). After recrystallization the crude aryl iodides, ArI, gave the four purified monoiodinated products in excellent yields (83-93%).

We have previously observed [8, 11] during our numerous experiments that in the presence of **water** (which always considerably diminishes the activity of all electrophilic iodinating species, owing to its high hydration power), the assumed transient *iodine(III)* species (briefly denoted above as I^{3+}) are *unstable* and very quickly vanish to form some *less reactive* and more stable *hydrated species*, probably I-O⁺ H₂. Presumably, these transformations would take place as follows [17]:

$$\begin{split} I_2(SO_4)_3 + 2H_2O &\to I_2(SO_4) + 2H_2SO_5 \\ I(SO_4)OSO_3H + H_2O &\to IOSO_3H + H_2SO_5 \\ I(OSO_3H)_3 + H_2O &\to IOSO_3H + H_2SO_5 + H_2SO_4 \\ I_2(SO_4) + 2H_2O + H_2SO_4 &\to 2I\text{-}O^+H_2 + 2HSO_4^- \\ IOSO_3H + H_2O &\to I\text{-}O^+H_2 + HSO_4^- \\ H_2SO_5 + H_2O &\to H_2SO_4 + H_2O_2 \end{split}$$

Hence, the *anhydrous* and *strongly acidic* conditions are indispensable to attain the possible highest yields of the assumed *iodine(III)* intermediates, ArISO₄, derived from the reacted *deactivated* arenes by their electrophilic substitution with I^{3+} ; cf. Eqs. 3, 5, 7, 9, and 10.

The characteristic feature of all the above oxidative iodination reactions (undergoing in agreement with Eqs. 8-10) is that diiodine is oxidized there by HIO₄ or HIO₃ to form some reactive transient species, I^+ or I^{3+} , whereas the both oxidants applied are reduced by diiodine to form *the same* transient species, I^+ or I^{3+} - which next react, more or less readily, with aromatic compounds, Ar-H, to form either ArI or ArISO₄ intermediates, respectively. Hence, no *strongly toxic* residues remain after the oxidative iodination reactions using the inorganic iodine(VII) or iodine(V) oxidants. Such iodination reactions are, indeed, *environmentally benign*.

In our opinion, particularly interesting are our iodination results obtained with *deactivated* arenes (Eq. 9 and 10). It has been a generalized opinion [19b] that inorganic iodine(VII) and iodine(V) oxidants are **mild** - hence they can hardly be appropriate for the effective oxidative iodination of *deactivated* aromatic substrates. Our experimental results with *deactivated* arenes clearly show that this notion is incorrect provided that *acidic* and *anhydrous* conditions are maintained in the reactions as shown in Eqs. 9 and 10.

Note. Quite recently, Russian chemists [49] have developed easy and effective procedures for iodination of *deactivated* aromatics, without casual oxidation of CH₃ or even CHO groups (e.g. they iodinated benzaldehyde in 61% yield). The reactions were carried out in conc. (90%) H₂SO₄ with some superactive iodine reagent "I⁺" prepared on a base of toxic iodine chloride with silver sulfate, at 0-20°C and within 15-150 min. Also, nearly all the former iodinating procedures suitable for

deactivated aromatics are briefly reviewed therein - with excluding, however, our two former papers [8, 11].

2.7. Aromatic Iodides from Activated Aromatics Obtained by an Improved, Acid- catalyzed Iodinating Procedure with (Diacetoxyiodo)benzene as Oxidant [15]

In a full-text version of our paper [15] there is a comprehensive **review** on the aromatic halogenation reactions with organic *trivalent* iodine reagents as the oxidants. So far, **nine** organic *iodine(III)* compounds have been used as the oxidants for the *oxidative halogenation* of aromatic compounds:



None of these reagents, however, permit *aromatic fluorination*. In view of electron density on the aromatic ring, it seems that halogenation of *chlorobenzene* represents the limit of the scope of the reactions with reagents 1-3 and 5-9; *more deactivated* aromatics cannot be halogenated with these reagents. But the evident advantages of these *solid* reagents are: 1) low toxity, 2) easy handling and simple experimental procedures, 3) usually high yields of halogenation products, 4) possibility (for reagents 1, 2, 5, 6 and 8) of recovery from the final reaction mixtures of *iodobenzene* which may be recycled after its oxidative conversion into the initial *iodine(III)* reagents, following known procedures [10, 21b]. Similarly, the polymeric reagent 3 can be recovered in good yield by filtration as poly(iodostyrene), and 3 can easily be regenerated and reused [50]. However, all the hypervalent iodine reagents are more or less light- and heat-sensitive, while 4 and 5 are also moisture-sensitive.

Reagents 2, 5 and 6 are commercially available with 2 being *notably cheaper* [48]. Moreover, pure 2 was synthesized by us in 79% yield with a novel procedure developed in our laboratory [10] (see Section 4.1); this procedure is ca. 5 times cheaper and 8-16 times faster than the former method [51]. Hence, in our work [15] we *considerably improved* the reported reaction conditions for the iodination of several *activated* arenes: benzene, iodobenzene, three xylenes, mesitylene, durene, di- and triphenylmethane, fluorene, fluoren-9-one, dibenzofuran, xanth-9-one, and uracil — using just the

reagent 2, which is more stable, preparatively more accessible, and notably cheaper than the moisturesensitive reagent 5, strongly recommended by other authors [19b] in place of 2.

Ogata and Aoki [52] first iodinated 1,3-xylene to 4-iodo-1,3-xylene (96%) in warm (60°C) acetic acid with reagent **2** as the oxidant, with iodobenzene, produced as the side product, being recovered. Next [19b], several *activated* arenes were iodinated in warm (50°C) acetic acid with $I_2/2$ to give mono-iodinated products, *within 1-5 hours*, in 56-89% yields. Togo and co-workers [50] have compared the reagent **2** and the polymer-supported reagent **3** in the oxidative iodinated nine activated arenes. With $I_2/2$ (in ethyl acetate, at room temperature or at 60°C) they monoiodinated nine activated arenes in 4-99% yields and *within 4-16 hours*; they also diiodinated mesitylene (96%), biphenyl (87%) and diphenyl ether (70%) at 60°C *after 16 hours*. After 1980, mono-, di-, and triiodinated *activated* arenes /hetero-arenes have instead been prepared with $I_2/5$ (in chlorinated solvents) in good or excellent yields, mainly at room temperature and within 0.25-2 hours [19b]. The other hypervalent iodine reagents listed above have, so far, been used to a lesser extent.

Transient acetyl hypoiodite (CH₃COI) was suggested as the iodinating species when reagent 2 is applied as the oxidant to iodinate aromatics [19b]. In the presence of strong acids, the reactivity of 2 is enhanced, because of its dissociation to more reactive cationic species [21b]:

$$PhI(OAc)_2 + H^{\textcircled{D}} \longrightarrow PhI^{\textcircled{D}}OAc + AcOH$$

As in our earlier works [8, 11], we deliberately applied *anhydrous* iodinating conditions, using AcOH/Ac₂O mixtures as the cheap solvents of choice [15]. The addition of varying *catalytic amounts* of conc. (98%) H₂SO₄ (from one drop up to 1 ml, depending on the reactivity of iodinated aromatics) *considerably accelerated* the iodination reactions studied; only fluorene was effectively iodinated in absence of H₂SO₄ at room temperature and within 15 minutes. We put forward the following reaction mechanism and stoichiometry [15]:

(1) PhI(OAc)₂ + I₂
$$\xrightarrow{H^{\oplus}}$$
 PhI + 2 AcOI (*transient acetyl hypoiodite*).
(2) 2 Ar-H + 2 AcOI \longrightarrow 2 ArI + 2 AcOH, hence:
(3) 2 Ar-H + I₂ + PhI(OAc)₂ \longrightarrow
AcOH/Ac₂O
 $\xrightarrow{H^{\oplus}; r.t.; 0.25 h}$ 2 Ar-I + PhI + 2 AcOH (Eq. 11).

The overall iodination rate is, in our opinion, mainly restricted by the rate of the first step (1), and is considerably accelerated by the *anhydrous* conditions applied as well as by *catalytic amounts* of conc. (98%) H_2SO_4 added to the iodinating (or brominating) reaction mixtures. The reactions proceeded quickly until the coloration of iodine faded, then the reaction mixtures were poured into excess aq.

 Na_2SO_3 solutions. The isolated crude products were purified by common methods. From the filtrates the iodobenzene was recovered in 40-75% yields.

Using our *improved* iodination procedure [15] it was possible to mono-, di-, or even triiodinate the studied aromatics, *at or near room temperature* and *within at most 15 minutes*, in 40-82% yields, except for uracil, which was also monoiodinated in 82% yield within 15 minutes, but at 40°C. For comparison, the *oxidative bromination* of mesitylene, durene and fluoren-9-one, carried out in a similar *anhydrous* system: arene/Br₂/2/AcOH/Ac₂O with only one drop of conc. H₂SO₄ added, resulted in tribromomesitylene (65%), dibromodurene (62%) and 2,7-dibromofluoren-9-one (73%). These bromination reactions even proceeded *much faster* than the corresponding iodination reactions. Hence, we recommend our *improved method* [15] as a very easy, fast, inexpensive and effective method to iodinate/brominate *activated* aromatic compounds.

Note. Dr. Maria Niemyjska [53] from our laboratory has carried out the oxidative iodination reactions using reagent **8** as the oxidant. The reactions took place in hot (95-100°C) AcOH/Ac₂O mixtures acidified with conc. H₂SO₄, within several hours. She thus obtained 4-iodoanisole (68%, admixed with its *ortho*-isomer) and 1-iodo-2-methoxynaphthalene (10%, admixed with 6-iodo-2-methoxynaphthalene), but she *failed* to iodinate acetanilide, 4-methoxybenzophenone, methyl salicylate, and mesitylene. These reactions were undertaken only for the sake of comparing **8** with the other hypervalent iodine reagents, and were unworthy of further study.

2.8. Aromatic Iodides from Deactivated Aromatics with a Stable Urea•H₂O₂ Complex as Oxidant [98]

Recently [98], we succeeded in oxidatively iodinating several *deactivated* arenes (in 40-88% yields), with a stable, commercially available **urea**• H_2O_2 complex used as the oxidant, viz.

$$I_{2} + 3H_{2}O_{2} \rightarrow [I_{2}O_{3}] + 3H_{2}O (a \text{ preliminary stoichiometry})$$

$$[I_{2}O_{3}] + 3H_{2}SO_{4} \rightarrow I_{2}(SO_{4})_{3} + 3H_{2}O$$

$$\frac{2Ar-H + I_{2}(SO_{4})_{3} \rightarrow 2ArISO_{4} + H_{2}SO_{4}, \text{ hence:}}{2 \text{ Ar-H} + I_{2} + 3 \text{ [urea]} \cdot H_{2}O_{2} + 2 H_{2}SO_{4} + 6 \text{ Ac}_{2}O \longrightarrow$$

$$AcOH/Ac_{2}O$$

$$\frac{40 \text{ oC}; 2 \text{ h}}{2 \text{ ArISO}_{4} (not \text{ isolated}) + 3 \text{ [urea]} + 12 \text{ AcOH} (Eq. 11')$$

After pouring the resulting (final) reaction mixtures into excess aq. Na₂SO₃ solutions (see Eq. 4 in Section 2.3), the isolated crude ArI were purified. It should be emphasized that this effective iodination method is evidently *environmentally benign*.

3. Syntheses of (Dichloroiodo)arenes

1346

(Dichloroiodo)arenes, ArICl₂, have found growing importance in modern organic synthesis [21]. More stable, solid ArICl₂, e. g. (*dichloroiodo)benzene*, are used as potent and fairly selective chlorinating and/or oxidizing agents. They have a practical advantage over elemental chlorine (dichlorine), due to their easy and safe handling. Moreover, they may be readily converted to other important organic hypervalent iodine reagents playing also an important role in organic synthesis, viz. iodosylarenes, iodylarenes, (diacyloxyiodo)arenes, (difluoroiodo)arenes [21], diaryliodonium salts [13], etc.

ArICl₂, yellow crystalline compounds, are light- and heat-sensitive and often unstable to storage. They do not usually give satisfactory microanalyses and, due to their thermal lability, their melting points are rather *uncertain*, depending upon the purity of the crude products prepared, the time elapsed since their preparation, and the rate of heating. Keefer and Andrews [54] have observed that in the cases of $4-O_2NC_6H_4ICl_2$ and $4-HOOCC_6H_4ICl_2$ all the dichlorine was expelled when their melting occured; the recorded melting points were the same as those of the initial iodoarenes, i. e. $4-O_2NC_6H_4I$ and $4-HOOCC_6H_4I$.

In 1885 Willgerodt [55] had prepared the first stable organic *iodine(III)* compound, (dichloroiodo)benzene: PhI + ICl₃ \rightarrow **PhICl₂** + ICl. ArICl₂ can also be made as follows [20]: Ar₂Hg + ICl₃ \rightarrow **ArICl₂** + ArHgCl; this reaction is still of preparative significance in the case of the vinyl-type organomercurials.

In 1886 Willgerodt [56] had developed the most common method up to now for preparing ArICl₂, by passing the stream of Cl₂ through solutions of ArI dissolved in CHCl₃, at 0 °C. The yields are generally excellent when this method is applied; ArICl₂ is not formed from ArI substituted with such groups as OH, NH₂, ethylenic double bonds, etc. Anhydrous conditions previously demanded [57] are not necessary [58]. Quite recently [59], the repeated preparations of PhICl₂ (in 94% crude yield) on a 20 kg scale have been conducted by the direct chlorination of PhI (dissolved in CH₂Cl₂), at -3 to +4 °C; next, it was possible to monochlorinate 4-aminoacetophenone with PhICl₂ on 24.8 kg scale, in 87% yield.

The *inconvenient* use of gaseous Cl₂ to afford ArICl₂ from ArI may be avoided as follows:

(i) by the action of hydrochloric acid on either iodosylarenes, ArIO, or (diacyloxyiodo)arenes, ArI(OAc)₂, previously otherwise obtained from ArI [21];

(ii) by using liquid SO_2Cl_2 added to a solution of ArI either in *wet* diethyl ether [60] or (much better) in 98% acetic acid [61]. Karele and Neiland [61] failed to chlorinate with SO_2Cl_2 three isomeric iodonitrobenzenes, 4-iodobenzoic acid, and 2-iodoacetophenone; for ten appropriate ArI the crude yields for ArICl₂ were 73-96%;

(iii) by the action of a liquid mixture: cobalt(III) acetate - KCl - 67% aq. CF_3COOH on PhI [62]; only iodobenzene was chlorinated there to give PhICl₂ in 75% crude yield;

(iv) by the use of hydrochloric acid oxidized *in situ* with sodium perborate tetrahydrate in either acetonitrile or CCl_4 containing a dissolved ArI [63]; when CCl_4 was applied as co-solvent, then the

chlorinations of three isomeric iodoanisoles were, in fact, *biphasic* ones. McKillop et al. [63] failed to chlorinate oxidatively 2-iodonitrobenzene; for fifteen appropriate iodobenzenes, including also 3- and 4-iodonitrobenzene, the obtained crude yields for ArICl₂ were 60-98%.

3.1. Biphasic Chlorination of Iodoarenes to (Dichloroiodo)arenes [12,18a, 64]

In 1991 we already reported [64] a *two-phase* (CCl₄/conc. hydrochloric acid) chlorination of iodobenzene and 4-iodoanisole to PhICl₂ and 4-MeOC₆H₄ICl₂, respectively, in yields exceeding 90%. The same *biphasic procedure* has later been applied in our laboratory [12] to chlorinate seventeen ArI compounds to the corresponding ArICl₂ in 50-98% crude yields, including also three isomeric iodonitro-benzenes, 3-HOOCC₆H₄I, 3-MeOOCC₆H₄I, and 3-iodobenzophenone.

The essence of this method is as follows. A solution of ArI in CCl_4 was vigorously stirred with excess conc. hydrochloric acid, at 0-5°C, and the powdered inorganic oxidant (KClO₃ or chlorinated lime) was very slowly added portionwise. Dichlorine was produced in the aqueous phase as follows:

KClO₃ (solid) + 6HCl (conc. aq.) $\rightarrow \uparrow 3Cl_2 + KCl + 3H_2O$, alternatively: CaCl(OCl) + 2HCl (conc. aq.) $\rightarrow \uparrow Cl_2 + CaCl_2 + H_2O$ (*less effective*)

Most of the dichlorine evolved was absorbed and consumed in the organic phase to chlorinate effectively a iodoarene dissolved therein. After 3 hours of stirring at $0-5^{\circ}$ C, the yellow precipitates, ArICl₂, were washed on the filter with dilute hydrochloric acid, water and CCl₄, and air-dried *in the dark*.

In our next work [18a] we used sodium peroxodisulfate (sodium persulfate), $Na_2S_2O_8$, a very potent, inexpensive [48] and often applied oxidant [40] to oxidize *in situ* conc. hydrochloric acid to chlorinate ArI to the corresponding ArICl₂, using either the *liquid-phase* (see Section 3.2) or *biphasic* chlorination procedures. Dichlorine was produced there as follows:

 $Na_2S_2O_8 + 2HCl \text{ (conc. aq.)} \rightarrow \uparrow Cl_2 + 2NaHSO_4$

However, a short preheating period (at $45-50^{\circ}$ C) was necessary to start the above oxidative reaction. Only PhI, 4-MeCONHC₆H₄I, 4-FC₆H₄I, 4-MeOC₆H₄I, and 2-IC₆H₄COOH were chlorinated by the *biphasic* method to the corresponding ArICl₂ obtained in 89-100% crude yields. 4-Acetamidoiodobenzene gave 4-acetamido(dichloroiodo)benzene in only 60% crude yield when it was chlorinated with the *liquid-phase* method, but its chlorination with the *biphasic* method afforded the same product in 90% crude yield. This example shows that for iodoarenes substituted with *hydrolyzable* groups, the *biphasic* chlorination is evidently preferable.

In comparison with the classic Willgerodt method [56-59], the *biphasic* chlorination method avoids the inconvenient use of gaseous Cl_2 , hence it is *relatively safe*. If an oxidant, e.g. KClO₃ [12], is very

slowly added to the vigorously stirred biphasic reaction mixture, then the escape of Cl_2 to the outer atmosphere is considerably limited. The limits of the scope of the two methods are the same or nearly so.

3.2. Oxidative Liquid-Phase Chlorination of Iodoarenes to (Dichloroiodo)arenes [14,18a,b]

In 1999 we reported [14] a novel, very simple, and efficient laboratory method for the conversion of seventeen iodoarenes to the corresponding (dichloroiodo)arenes, with including 2- and 3-iodonitrobenzene and 3-iodobenzoic acid; although *we failed* to chlorinate 4-iodonitrobenzene, 4-iodobenzonitrile, and 1, 3, 5-trichloro-2-iodobenzene. We used conc. hydrochloric acid oxidized *in situ* with varied amounts of **chromium(VI) oxide**, a common and easily handled oxidant [37, 40], dissolved in aq. acetic acid, and acting upon a dissolved (or suspended) iodoarene. The reactions proceeded according to the following stoichiometry:

The crude yellow products precipitated by water were washed well on the filter with ice-cold water to remove $CrCl_3$, HCl, and AcOH, then with CCl_4 to remove most of the unreacted ArI, and **air-dried** *in the dark*; we observed that by drying the crude $ArICl_2$ in a vacuum desiccator, the chlorine percentage was *lowered*: $ArICl_2 \rightarrow ArI + \uparrow Cl_2$.

Note: *easily oxidizable* ArI, e.g. 4-iodoanisole and iodotoluenes, should be added to the reaction mixture **after** the addition of the whole portion of conc. HCl. Otherwise, the final yields of the corresponding $ArICl_2$ were very low.

The same procedure is effective for the aromatic chlorination of *activated arenes*. Exemplarily, we obtained the purified chlorinated products: 2-chloro-4-nitroanisole (64%), 2-chloro-4-nitroacetanilide (58%), and 3,5-dichloro-4-methoxybenzoic acid (42%) from the respective arenes.

Later, we reported [18a] the preparation of (dichloroiodo)arenes, $ArICl_2$, from the respective iodoarenes, ArI, by the use of hydrochloric acid oxidized *in situ* with **sodium peroxodisulfate** (sodium persulfate), $Na_2S_2O_8$, in acetic acid containing a dissolved ArI; it was necessary to warm up preliminarily the stirred reaction mixture to 45-50°C to initiate the oxidative chlorination of ArI. *We failed* to chlorinate 2-iodo-1,4-xylene, iodomesitylene, and 1,3,5-trichloro-2-iodobenzene. For sixteen appropriate ArI, with including *all* the isomeric iodonitrobenzenes and iodobenzoic acids, the crude yields for $ArICl_2$ were 60-100%. The chlorination reactions underwent according to the following stoichiometry:

ArI + Na₂S₂O₈ + 2 HCl (conc. aq.)
$$\longrightarrow$$

aq. AcOH
$$\underbrace{45-50 \text{ °C, then r.t.; 4 h}}_{60-100\% (crude yields)} \text{ ArICl}_2 + 2 \text{ NaHSO4}$$
(Eq. 13).

Finally, we presented [18b] one improved (cf. Ref. 63) and eight novel oxidative, *liquid-phase* chlorination procedures for the preparation of (dichloroiodo)arenes, ArICl₂, from always the same *nine* exemplary iodoarenes, ArI, using KMnO₄, activated (85%) MnO₂, KClO₃, NaIO₄, NaIO₃•H₂O, conc. nitric acid, NaBO₃•H₂O, 2Na₂CO₃•3H₂O₂, and a stable urea•H₂O₂ complex, used as the oxidants, which oxidized hydrochloric acid to produce *in situ* an active chlorine (*in statu nascendi*). The crude yields for the nine ArICl₂ obtained were good or excellent (63-99%).

The following general reaction stoichiometries (Eqs. 14-22) may be deduced for the oxidative, *liquid-phase* chlorination procedures applied in our work [18b], viz.

$$ArI + NaBO_3 \bullet H_2O + 2HCI \rightarrow ArICl_2 + [NaBO_2] + H_2O$$
(Eq.14)

McKillop et al. [63] applied NaBO₃•4H₂O as the oxidant used in the 400% excess, using acetonitrile (or, only for iodoanisoles, carbon tetrachloride) as co-solvent. They stirred the reaction mixtures, at room temperature, mostly for 2 hours, but for 1,4-diiodobenzene and 4-iodonitrobenzene the stirring had to be prolonged *to 3 days*; no chlorination of 2-iodonitrobenzene was attained even after 3 days. In our present work [18b], Na₂BO₃•H₂O was used in only the 200% excess, the reaction times did not exceed 2 hours, but we unexpectedly *failed* to chlorinate *4-iodonitrobenzene*, in spite of several attempts (we also met with the same failure using the other oxidants in our work [18b]).

$$ArI + HNO_3 + 3HCl \rightarrow ArICl_2 + [NOCl] + 2H_2O$$
(Eq. 15)

We *excluded* the possibility of chlorination of ArI with NOCl (formed there as a side product [65]), carrying out the separate, *biphasic* experiments. ArI were dissolved in CHCl₃ or CCl₄, excess conc. hydrochloric acid was added, then solid NaNO₂ was slowly added portionwise with vigorous stirring. In the aqueous phase the following reaction took place [66]: NaNO₂ + 2HCl \rightarrow NOCl + NaCl + H₂O. The NOCl formed was surely present in excess in the organic phase, since it is readily soluble in chlorinated aliphatic hydrocarbons. However, *no trace* of any yellow coloration was developed, characteristic of ArICl₂.

$$5\text{ArI} + 2\text{KMnO}_4 + 16\text{HCl} \rightarrow 5\text{ArICl}_2 + 2\text{MnCl}_2 + 2\text{KCl} + 8\text{H}_2\text{O}$$
(Eq. 16)
ArI + activated MnO₂ + 4HCl \rightarrow ArICl₂ + MnCl₂ + 2H₂O (Eq. 17)

We observed that $ArICl_2$, produced according to Eqs. 16 and 17, were *more contaminated* than those obtained with the other procedures.

$$3ArI + NaIO_4 + 8HCl \rightarrow 3ArICl_2 + [ICl] + NaCl + 4H_2O$$
(Eq. 18)
$$2ArI + NaIO_3 \bullet H_2O + 6HCl \rightarrow 2ArICl_2 + [ICl] + NaCl + 4H_2O$$
(Eq. 19)

So far, we have had no experimental evidence suggesting that iodine chloride was formed as a side product in the reaction mixtures studied. Nevertheless, high crude yields of $ArICl_2$ formed from ArI in the chlorination reactions undergoing in agreement with such the supposed stoichiometries would suggest that Eqs. 18 and 19 are plausible.

$$3ArI + KClO_3 + 6HCl \rightarrow 3ArICl_2 + KCl + 3H_2O$$
(Eq. 20)

Knowing the explosive properties of KClO₃, it **must not** be pulverized in a mortar prior to its use in the reactions.

$$3\text{ArI} + 2\text{Na}_2\text{CO}_3 \bullet 3\text{H}_2\text{O}_2 + 10\text{HCl} \rightarrow 3\text{ArICl}_2 + 4\text{NaCl} + 2\text{CO}_2 + 8\text{H}_2\text{O}$$
(Eq. 21)
ArI + urea • H₂O₂ + 2HCl \rightarrow ArICl₂ + [urea] + 2H₂O (Eq. 22)

Sodium percarbonate and sodium perborate are both available at a low price, and are widely used in chemical laboratories or in industry [67]. They may be considered as "dry carries" of the hazardous and unstable hydrogen peroxide, are easy to handle, safe and stable at room temperature; the same is true in respect to a commercially available, *stable* complex of urea with hydrogen peroxide [68]. Their ability to release oxidative species in organic media has made them useful reagents in organic synthesis. Hence, they were also checked in our work [18b].

When the oxidative chlorination reactions are carried out in a *liquid phase* containing ArI, an appropriate strong oxidant, and conc. hydrochloric acid mixed with an inert solvent, then a very active chlorine, *in statu nascendi*, is instantly produced there, which is apt to react not only with the iodine atoms of ArI, but also with aromatic rings of highly activated ArI, e. g. iodoanisoles, which were formerly chlorinated by McKillop with co-workers [63]. There underwent rapid aromatic chlorinations when acetonitrile was used as co-solvent (*a liquid-phase protocol*). 3-Iodoanisole gave 2,4,6-trichloro-3-methoxyiodobenzene, whereas 4-iodoanisole gave 3-chloro-4-methoxy(dichloroiodo)benzene initially, but this rapidly decomposed to give 3,5-dichloro-4-methoxyiodobenzene. However, the same iodoanisoles could be satisfactorily chlorinated to the corresponding ArICl₂ using CCl₄ as co-solvent (*a biphasic protocol*) - it means that the *two-phase* systems allow *milder* chlorination conditions. The latter may be explained as follows. An active chlorine *in statu nascendi*, formed in the aqueous phase of the biphasic system, is very quickly deactivated there to form the "ordinary" dichlorine, Cl₂, which next is readily absorbed and consumed in the organic phase containing ArI to produce there ArICl₂. Similarly, in the Willgerodt method [56-59], the stream of this "ordinary" dichlorine, Cl₂, is passed

through solutions of ArI in inert solvents to form there ArICl₂. Thus, we previously expressed the opinion [12] that "the limits of the scope of the two methods are the same or nearly so." Later [18a], we concluded our chlorination experiments as follows: "... it is preferable to obtain ArICl₂ from iodoanisoles and highly activated ArI with using either the biphasic method or the classic Willgerodt method, where such ArI and a *less active* molecular chlorine Cl₂, are both dissolved and reacting in an inert solvent. Contrariwise, more suitable for *deactivated* ArI seem to be appropriate liquid-phase chlorinating protocols - because such ArI are reacted upon with some *more active*, though short-living, chlorine species, formed there *in statu nascendi*."

3.3. A One-Pot Method for Preparing (Dichloroiodo)arenes from Arenes [18c]

Almost all former methods of preparing ArICl₂ demanded the use of *costly* **iodoarenes**, ArI, as the starting substrates to be then chlorinated at their iodine atoms. In our newest paper [18c] we have presented a quite novel, **one-pot** (two-stage) method for preparing eleven exemplary ArICl₂ from the respective **arenes**, Ar-H, used as the starting substrates to afford ArICl₂, isolated in 46-88% crude yields, which were *possibly optimized*. Ar-H were, at first, oxidatively substituted in appropriate, *anhydrous* I₂/**NaIO**₄ *or* **NaIO**₃/AcOH/Ac₂O/conc. H₂SO₄ mixtures (discussed previously in Section 2.6), with some transient *iodine(III)* species, denoted as I^{3+} , to form *in situ* the respective organic *iodine(III)* intermediates, denoted as ArISO₄. Formerly [17], the resulting reaction mixtures, containing ArISO₄ intermediates, were poured into excess aq. Na₂SO₃ solutions to afford *iodoarenes*, ArI (see Eq. 4) In our work [18c], ArICl₂ were precipitated from the same, resulting (final) reaction mixtures, containing ArISO₄ intermediates, by adding to them a large excess of conc. hydrochloric acid. Thus, the essence of this quite novel method is following:

1) $2I_2 + 3I(VII) \rightarrow 7I^{3+}$ or $I_2 + 3I(V) \rightarrow 5I^{3+}$ (preliminary stoichiometries) (these reactions undergo in anhydrous, acidic solutions - see Section 2.6)

2) Ar-H +
$$\mathbf{I}^{3\oplus}$$
 $\xrightarrow{-H^{\oplus}}$ ArI $^{2\oplus}$ $\xrightarrow{+H_2SO_4}$ ArISO₄ (not isolated).
3) ArISO₄ (not isolated) + 2HCl (conc. aq.) $\rightarrow \downarrow$ ArICl₂ + H₂SO₄ (Eq. 23)

The yellow precipitates were collected by filtration, washed well with ice-cold water, a little CCl₄, and air-dried *in the dark*.

When **NaIO**₄ was used as the oxidant, the following $RC_6H_4ICl_2$ were obtained: R = H (88%), 4-F (70%), 4-Cl (70%), 4-Br (75%), 4-I (81%), 3-COOH (86%), 3-COOMe (69%), 3-COOEt (74%), 3-CF₃ (57%), 4-OMe (57%); also 2,4-Cl₂C₆H₃ICl₂ (46%) was obtained from 1,3-C₆H₄Cl₂. When **NaIO**₃ was used as the oxidant, the following $RC_6H_4ICl_2$ were obtained: R = H (87%), 4-Br (63%), 3-COOH (80%), 3-COOMe (64%). Of course, only those isomeric $RC_6H_4ICl_2$ may predominantly be obtained

from the monosubstituted benzenes, RC_6H_5 , which are formed in agreement with common orientation rules in the electrophilic substitutions of RC_6H_5 with *strongly electrophilic* I^{3+} transient species. We *failed* to synthesize ArICl₂ from toluene, phenetole, 4-nitroanisole, 4-nitrotoluene, 2methylbenzaldehyde, 2-methoxybenzaldehyde, 4-chlorobenzaldehyde, 4-toluic acid, and 4bromobenzoic acid. Further studies are necessary to establish the limits of the scope of this novel method, which is evidently *environmentally benign* - hence it would be particularly suitable for *largescale* preparations of ArICl₂, e.g. PhICl₂ (cf. Ref. 59).

4. Syntheses of (Diacetoxyiodo)arenes

(Diacetoxyiodo)arenes, ArI(OAc)₂, and particularly the parent (*diacetoxyiodo*)benzene, PhI(OAc)₂, have been known for a long time [21, 56]. They are potent, often fairly selective, oxidizing agents, hence the interest in ArI(OAc)₂ and PhI(OAc)₂ is growing rapidly, as demonstrated by a number of recent books [21] and reviews [69]. They are also used for the facile syntheses of, for example, [*bis*(trifluoroacetoxy)iodo]arenes, [hydroxy(tosyloxy)iodo]arenes (*selective oxidants*), aromatic iodonium salts (*arylating reagents*), etc. [21, 69].

There are several preparative methods for these compounds. So far the *substrates* have generally been [21, 56, 69]:

(i) *iodosylarenes* dissolved in glacial acetic acid;

(ii) (*dichloroiodo*)*arenes* in which the chlorine atoms are exchanged by acetoxy groups coming either from silver, lead(II) or sodium acetate, or from acetic acid in the presence of mercury(II) oxide in chlorinated solvents [61];

(iii) *iodoarenes* are oxidized in *warm* glacial acetic acid by either peracetic acid, or sodium perborate [51], or electrolytically.

The standard, and most general, method for the synthesis of $ArI(OAc)_2$ (oxidative diacetoxylation of ArI by warm peracetic acid) is, in fact, a very prolonged reaction (12-16 hours), and the utmost care should be taken to maintain the exact temperature, 40°C [21b]. $ArI(OAc)_2$ are generally crystalline compounds, fairly stable in the air, which may be stored for long periods by avoidance of light, and preferably in a cooler.

4.1. A Two-Step Conversion of Iodoarenes to (Diacetoxyiodo)arenes with Chromium(VI) Oxide as Oxidant [10]

In our earlier papers [3-5, 6b, 9] many **short-cut** syntheses of diaryliodonium salts were reported (for more details see Section 6). We oxidized various ArI (*excluding* those substituted solely with stronger electron-donating groups, e.g. OMe, NHAc) with appropriate mixtures of *anhydrous* CrO_3 /AcOH/Ac₂O/conc. H₂SO₄, immediately followed by the acidic coupling of the *in situ* formed *iodine*

(*III*) intermediates, ArISO₄ and/or ArI(OSO₃H)₂, with many activated arenes, Ar'-H. The soluble diaryliodonium hydrogensulfates thus obtained, Ar(Ar') Γ^{+} HSO₄⁻, were next precipitated out in the form of *insoluble* diaryliodonium bromides, iodides, or perchlorates. Alternatively, by pouring the deep-green solutions containing Cr(III) salts and ArISO₄ and/or ArI(OSO₃H)₂ into excess aq. Na₂SO₃ solutions, the respective ArI were obtained in high yields; see Eqs. 3 and 4 in Section 2.3.

In our work [10] we applied the same method to the oxidation of seventeen ArI with the appropriate $CrO_3/AcOH/Ac_2O/conc$. H_2SO_4 liquid systems, followed by mixing the resulting deep-green reaction mixtures with excess 20% aq. ammonium acetate solutions. Crude crystalline ArI(OAc)₂, obtained according to Eqs. 24 and 25, viz.

$$3 \operatorname{ArI} + 2 \operatorname{CrO}_3 + 6 \operatorname{Ac}_2 O + 6 \operatorname{H}_2 \operatorname{SO}_4 \longrightarrow$$

$$40 \operatorname{^{o}C}, 30 \operatorname{min} \longrightarrow 3 \operatorname{ArISO}_4 (not isolated) + \operatorname{Cr}_2(\operatorname{SO}_4)_3 + 12 \operatorname{AcOH}$$
(Eq. 24).

 $3 \text{ ArISO}_4 (not isolated) + 6 \text{ AcONH}_4 (aq.) \longrightarrow$

$$\frac{5 \text{ oC; fast reaction}}{58-82\%} \oint 3 \text{ ArI(OAc)}_2 + 3 \text{ (NH4)}_2\text{SO}_4 \tag{Eq. 25}$$

were admixtured with a little of the *hydrolyzed* yellowish side products, viz. ArI(OH)OAc, ArIO, and [ArI(OAc)]₂O. Crude yellowish ArI(OAc)₂ were washed on the filter with a cold 10% aq. AcOH (in which they are *less soluble* than in water), air-dried in the dark, and recrystallized from either AcOH or AcOMe mixed with **Ac₂O** (9:1) to acetylate the yellowish side products back to ArI(OAc)₂. Then, either hexane or Et₂O were added in excess to the cooled solutions to improve the crystallization yields. After washing on the filter with hexane or Et₂O, the final yields of pure ArI(OAc) were 58-82%. This method is *unsuitable* for iodoanisoles and iodoacetanilides. 4-Iodotoluene gave 4-(diacetoxyiodo)toluene in only 20% yield; as a side product we identified 4-iodobenzoic acid (40%), the unreacted 4-IC₆H₄Me as well as 4-tolyl acetate by TLC. Pure Ph(OAc)₂ was obtained from PhI in 79% yield. Taking into account the reaction times as well as total amounts, and the respective costs, of all the reagents and solvents necessary to prepare pure ArI(OAc)₂, our method [10] is *8-16 times faster* (30 min as compared with 4-8 h) and *ca. 5 times less expensive* than the method of McKillop and Kemp [51].

4.2. (Diacetoxyiodo) arenes from Iodoarenes with Sodium Perborate Monohydrate as Oxidant [71]

McKillop and Kemp [51] oxidized thirteen ArI to the corresponding $ArI(OAc)_2$ with NaBO₃•4H₂O (900% excess) in a large volume of glacial AcOH within 4-8 hours, at 40-45°C. After recrystallizations from AcOH/hexane or cyclohexane, they obtained pure $ArI(OAc)_2$ in 66-80% yields; no attempt was made to optimize yields.

In our unpublished work [71] we used *acetic anhydride* as the solvent of choice and NaBO₃•H₂O as the oxidant, which allowed to lower considerably an excess of the oxidant applied to only 200%. The reactions underwent as follows:

ArI + NaBO₃ · H₂O + 2 Ac₂O
$$\longrightarrow$$

$$\frac{40 \text{ oC, } 2\text{-4 h}}{60\text{-}87\% (crude vields)} \text{ ArI(OAc)}_2 + [NaBO_2] + 2 \text{ AcOH}$$
(Eq. 26).

This work is still in progress and will be published soon.

4.3. (Diacetoxyiodo) arenes from Iodoarenes with Sodium Periodate as Oxidant [18e]

The essence of this novel method is following:

$$R \xrightarrow{I + \text{NaIO}_4 + \text{Ac}_2\text{O}} I + \text{NaIO}_4 + \text{Ac}_2\text{O} \xrightarrow{\text{AcOH/Ac}_2\text{O/AcONa}}_{45-77\%} R \xrightarrow{I(\text{OAc})_2 + \text{NaIO}_3}_{R}$$

R = H, 4-F, 4-Br, Cl (three isomers), Me (three isomers), OMe (three isomers).

The parent pure PhI(OAc)₂ was obtained from PhI in 73% yield; this method is *not applicable* for ArI substituted with strong electron-withdrawing groups. The presence of *sodium acetate* (in stochiometric quantities) in the reaction mixtures is indispensable - without its addition the oxidation reactions did *not* proceed. When sodium acetate was replaced for pyridine, then the final yields of ArI(OAc)₂ were lowered by ca. 10-20%. The purities and homogeneities of the purified ArI(OAc)₂ were checked by TLC, mixed melting points with authentic specimens as well as with ¹H-NMR spectra and elemental analyses. This novel method nicely *complements* our former method [10], which was hardly applicable for iodotoluenes and quite inappropriate for ArI substituted with stronger electron-donating groups, e. g. iodoanisoles and iodoacetanilides.

5. Syntheses of Iodylarenes

Rather few iodylarenes, ArIO₂, have been used as useful oxidants in organic synthesis. These include *iodylbenzene*, PhIO₂, and some of its substituted derivatives, e.g. 3-iodylbenzoic acid or 4-*tert*-butyliodylbenzene. In some cases, the reactivity of PhIO₂ was increased in the presence of CF₃CO₂H. But their former limited applications were partly due to their polymeric nature which makes them insoluble in most ordinary solvents, excepting *water*. Their synthetic utility is at present quickly growing and is more appreciated. The most applied reagents of *iodine(V)* are: cyclic "2-iodylbenzoic acid" (1-oxido-1-hydroxy-1,2-benziodoxol-3(1*H*)-one) and particularly the Dess-Martin reagent (1,1,1-

1355

triacetoxy-1,1-dihydro-1,2-benziodoxol-3(1*H*)-one), which synthetic applications, as versatile selective oxidants, are growing nearly exponentially; for more details see Refs. [21, 69a]. ArIO₂ are also useful in the preparation of iodonium salts, $Ar_2I^+X^-$ or $Ar(Ar')I^+X^-$ [56, 72, 73].

 $ArIO_2$ are *polymeric* and they not dissolve in ordinary organic solvents. They are rather stable thermally, but their melting points are, in fact, their decomposition points, accompanied by explosion. A violent decomposition of PhIO₂ (dry sample) has been induced by scraping with a spatula [21b].

Sharefkin and Saltzman [74] briefly summarized the former methods of preparing PhIO₂ as follows (see their Refs. 4-14): "PhIO₂ has been prepared by the disproportionation of PhIO, by oxidation of PhIO with hypochlorous acid or bleaching powder, and by oxidation of PhI with hypochlorous acid or with sodium hydroxide and bromine. Other oxidizing agents used with PhI include air, chlorine in pyridine, Caro's acid, conc. chloric acid, and peracetic acid solution. Hypochloric oxidation of PhICl₂ has also been employed". Their one-step method of preparing PhIO₂ (which seems general for ArIO₂, at least those with *electron-releasing* substituents) depended on the oxidation of PhI with commercial 40% peracetic acid, which resulted in pure PhIO₂ (72-80%). Indian chemists [75] also briefly reviewed the former methods of preparing ArIO₂ (see their Refs. 1-8), and made the following comments: "The reported methods of Lucas and Kennedy [76] for the preparation of PhIO₂ from PhI proved to be lenghty and irksome in our hands while that of Sharefkin and Saltzman [74] was neither economical nor free of danger for the use of peracetic acid, though the yield of the desired product in this single step reaction was excellent. Other effective oxidizing agents employed are oxygen, perbenzoic acid, potassium peroxymonosulfate,...and dibenzoyl peroxide and iodine;...also the preparation of ArIO₂ from ArI was reported using nitric acid and trifluoroacetic anhydride followed by hydrolysis of the intermediate PhI(OCOCF₃)₂", and next followed by steam-distillation. They developed the preparation of ArIO₂ by the oxidation of six ArI using KBrO₃ under acidic conditions, which afforded the following ArIO₂: PhIO₂ (45%), 4-BrC₆H₄IO₂(98%), 3-O₂NC₆H₄IO₂ (46%), 4-O₂NC₆H₄IO₂ (88%), 2-HOOCC₆H₄IO₂ (95%), and 3-MeC₆H₄IO₂ (poor yield). KBrO₃ and dilute H₂SO₄ produced PhIO₂ from PhIO in excellent yield.

The treatment of halobenzenes, benzoic acid, or nitrobenzene, dissolved in unspecified solvents, with HIO₃ and conc. H₂SO₄ *possibly* resulted in the formation of the respective iodylarenes, **ArIO**₂; next, they may be reduced to the corresponding iodides [77]. This short communication, with no experimental and physico-chemical support, was never followed up either then or after 1969 by the more detailed experimental paper. This should be checked and, possibly, extended, because this *direct method* seems to be promising.

A considerable progress in the synthesis of $ArIO_2$ is greatly due to a quickly growing importance of "2-iodylbenzoic acid" and its triacetoxy derivative, i.e. the Dess-Martin reagent (*vide supra*). From the last short review [78] it is seen that they were obtained, most satisfactorily, in the following ways [79, 80]:



The monoacetylated form of the above "2-iodylbenzoic acid", i.e. 1-acetoxy-1,2-benziodoxol-3(1H)-one-1-oxide, is probably the actual oxidizing species [81]. Note. "2-Iodylbenzoic acid" is explosive on heating above 200°C and also upon impact; the Dess-Martin reagent explodes violently on heating under confinement, at 130°C [21b].

5.1. Early Results: Biphasic Oxidation of Iodobenzene to Iodylbenzene [64]

In 1990/1991 a *biphasic* oxidation of PhI to PhIO₂ (77%), in the presence of a phase transfer catalyst, $Bu_4N^+Br^-$, was performed as follows [64]. PhI was dissolved in CCl₄, while excess NaOCl and a catalytic amount of $Bu_4N^+Br^-$ were dissolved or suspended in water. The both phases mixed together in the reaction vessel were vigorously stirred and warmed up to boiling under a reflux condenser. 800% excess conc. hydrochloric acid was slowly added dropwise to the boiling mixture within a few hours. After cooling, crude PhIO₂ was collected, washed with water, and recrystallized from boiling water. KMnO₄ used as the oxidant was somewhat less effective, and yielded crude PhIO₂ contaminated with MnO₂. Since Dr. J. Golinski left our laboratory, this work has not been continued.

In 1993 a simple *two-phase* method was published [82], involving the direct oxidation of ArI using aq. hypochlorite and phase transfer catalyst; for example [21b]:

PhI (in CH₂Cl₂)
$$\xrightarrow{\text{HOCl/H}_2O}$$
 PhIO₂ (59-81%).

5.2. Iodylarenes from Iodoarenes with Sodium Periodate as Oxidant [18e]

The *heterophasic* reactions between various iodoarenes suspended in boiling aq. $NaIO_4$ solutions proceeded smoothly within 8-16 hours to give the corresponding, colorless iodylarenes in 58-91% crude yields, viz.

$$R \xrightarrow{H_2O (reflux)} I + 2 \text{ NaIO4} \xrightarrow{8-16 \text{ h}} R \xrightarrow{IO_2 + 2 \text{ NaIO3}} (Eq. 27).$$

When the above reactions were completed, the cooled reaction mixtures were diluted with cold water. The collected precipitates were washed on the filter with cold water, next with CHCl₃ or acetone (to remove the unreacted ArI), and air-dried in the dark. The following crude $RC_6H_4IO_2$ were obtained (yields, %): R = H (86), 4-OMe (85), 2-Me (61), 3-Me (77), 4-Me (80), 4-F (91), 3-Cl (75), 4-Cl (80), 4-Br (73), 3-NO₂ (85), 4-NO₂ (58), and 3-COOH (89). Iodometric titrations [56, 97] showed that the contents of ArIO₂ in the crude products varied between 98.5 and 99.2%. Small samples of the crude products were recrystallized from boiling water to give the analytically pure specimens, which melting/detonation points were close to those reported in the literature. Their IR spectra displayed the characteristic frequencies for iodylarenes [21a].

The reaction mixtures discussed above were nearly *neutral*. When they were made *alkaline*, the unreactive $Na_3H_2IO_6$ precipitated out, and the reactions did *not* proceed. The same *negative* results were obtained, when **NaIO**₄ was replaced with **NaIO**₃ used as the oxidant under the same reaction conditions. When the reaction mixtures discussed above were *acidified* with either acetic acid or aq. H_2SO_4 , then the yellow colored *iodosylarenes*, ArIO, were the main products of the reactions; they were not studied further.

Exceptionally, when 2-iodobenzoic acid was oxidized as above with boiling aq. NaIO₄ solution, then "2-iodosylbenzoic acid" (i.e. 1-hydroxy-1,2-benziodoxol-3(1H)-one) was isolated in 76% crude yield. After recrystallization from boiling water, its structure was confirmed as above. Contrariwise, when *sodium salt* of 2-iodobenzoic acid was reacted as above with boiling aq. NaIO₄ solution, then this resulted in the formation of "2-iodylbenzoic acid" (i.e. 1-hydroxy-1,2-benziodoxol-3(1H)-one-1-oxide), isolated in 71% crude yield. After recrystallization from boiling water, its chemical structure was supported analytically, spectrally (IR), and with its characteristic melting/detonation point. Further informations will be given in our just completed paper [18e].

6. Syntheses of Diaryliodonium Salts

Symmetric and unsymmetric diaryliodonium salts, $Ar_2I^+X^-$ and $Ar(Ar')I^+X^-$, represent a very important class of aromatic *iodine(III)* compounds. They are widely used in organic synthesis as *arylating reagents* for organic and inorganic nucleophiles, some of them display *a biological activity* and *photochemical properties*, hence they are often applied as efficient catalysts for radiation initiated polymerization [21a,b]. Generally, their reactivity is less pronounced than that of other hypervalent iodine compounds. In several of their reactions relatively drastic conditions may be necessary, especially for the least reactive heterocyclic iodonium salts. The search for optimum conditions is often desirable even for well-established reactions, by applying new findings concerning the use of specific solvents, catalysts or radical traps [21b].

Diaryliodonium salts are generally solid compounds, mostly stable towards heat, oxygen and humidity; they are mildly light-sensitive and should be stored in the dark, without refrigeration [21b].

A large number of methods are available for the preparation of diaryliodonium salts. A brief account of the most often used methods is given by Varvoglis in his latest book [21b]; for more details see Refs. 21a, 69a, 83, 84.

6.1. Short-Cut Syntheses of Diaryliodonium Salts with Chromium(VI) Oxide as Oxidant [3-5, 6b, 9, 13]

In 1995 we reported [3] a one-pot (short-cut) synthesis of sparingly soluble diaryliodonium bromides from various ArI oxidized with anhydrous $CrO_3/AcOH/Ac_2O/conc$. H_2SO_4 mixtures, then coupled *in situ* with various arenes and, finally, precipitated with excess aq. KBr solutions. The essence of this novel method is shown below (for experimental details see [3]).

$$3 \operatorname{ArI} + 2 \operatorname{Cr}(\operatorname{VI}) \xrightarrow{H^{+}} 3 \operatorname{ArI}^{2+} + 2 \operatorname{Cr}(\operatorname{III}) (a \text{ preliminary stoichiometry})$$

$$3 \operatorname{ArI}^{2+} + 6 \operatorname{H}_{2}\operatorname{SO}_{4} \longrightarrow 3 \operatorname{ArI}(\operatorname{OSO}_{3}\operatorname{H})_{2} + 6 \operatorname{H}^{+}, \text{ hence:}$$

$$3 \operatorname{ArI} + 2 \operatorname{CrO}_{3} + 6 \operatorname{Ac}_{2}\operatorname{O} + 9 \operatorname{H}_{2}\operatorname{SO}_{4} \longrightarrow$$

$$\xrightarrow{\operatorname{AcOH}/\operatorname{Ac}_{2}\operatorname{O}} 3 \operatorname{ArI}(\operatorname{OSO}_{3}\operatorname{H})_{2} + \operatorname{Cr}_{2}(\operatorname{SO}_{4})_{3} + 12 \operatorname{AcOH} \qquad (Eq. 28).$$

$$4 \operatorname{-X-C}_{6}\operatorname{H}_{4}\operatorname{I} \xrightarrow{\operatorname{CrO}_{3}/\operatorname{AcOH}/\operatorname{Ac}_{2}\operatorname{O}/\operatorname{H}_{2}\operatorname{SO}_{4}}_{(\text{oxidation stage})} \longrightarrow \left[4 \operatorname{-X-C}_{6}\operatorname{H}_{4} \operatorname{-I}(\operatorname{OSO}_{3}\operatorname{H})_{2} \right] \longrightarrow (\operatorname{not isolated})}^{\operatorname{Ar}\operatorname{H}} \operatorname{CrO}_{4 \operatorname{KBr}} \operatorname{Ker}_{6} \operatorname{H}_{4} \operatorname{-I}_{6} \operatorname{Ker}_{6} \operatorname$$

For X = H, CH_3 , Cl, NO_2 ; if $X = OCH_3$ or $NHCOCH_3$, effective oxidation was *not* achieved.

Ar-H = benzene, toluene, mesitylene, anisole, chlorobenzene, acetanilide, anisic acid, and thiophene.

We have established that our one-pot method is *unsuitable* to couple the oxidized iodoarenes with nitrobenzene, benzoic acid and its methyl and ethyl esters, benzonitrile, acetophenone, benzophenone, cinnamic acid and its ethyl ester, *beta*-nitrostyrene, and chalcone under reaction conditions given in our paper [3].

The soluble diaryliodonium hydrogensulfates, $Ar(Ar')I^{+}HSO_{4}^{-}$, formed in the last stage were metathesized into the *insoluble bromides* by adding excess aq. KBr to the final, deep-green reaction mixtures, also containing the soluble chromium(III) salts. The chromium salts were completely *washed off* with water on the filter, followed with acetone (which removed organic impurities), leaving the nearly pure *diaryliodonium bromides* in 20 - 88% crude yields. The sparingly soluble diaryliodonium

iodides or *perchlorates* may also be obtained by the same one-pot method by adding excess aq. KI or HClO₄ to the final, deep-green reaction mixtures; they were obtained in 40 - 89% crude yields.

Note. Mgr Anna Kryska from our laboratory has recently established that CrO_3 (Eq. 28) may be replaced by *finely powdered* $K_2Cr_2O_7$, with preserving the same crude yields of the iodonium bromides/iodides.

There are repeated reports in the literature [84, 85] that diaryliodonium halides are *thermally unstable* on prolonged heating, hence their recrystallizations should be carried out *as quickly as possible* from boiling solvents. Analytical purity of the iodonium bromides and iodides obtained by us was achieved by *quick* dissolving small crude samples in boiling MeOH, followed by the immediate hot filtration neglecting the losses, and rapid cooling. More stable thermally iodonium perchlorates were recrystallized in usual way; *after drying* they should be handled with care.

The aforementioned new method [3] is easier and shorter than many earlier methods [21, 69a, 72, 83, 84], hence this method was also used in our subsequent work [4, 5, 6b, 9, 13] (see Sections 6.2 and 6.3).

In the same paper [3] and in the following paper [6b] we reported the *oxidative metatheses* in the *crude* iodonium bromides, iodides and chlorides into the corresponding *pure* tetrafluoroborates, tosylates, trifluoroacetates, etc. This will be dealt with in Section 7.1.

6.2. Iodonium/Diiodonium Salts Derived from Various Tricyclic Carbo- and Heteroaromatic Systems [4, 5, 9]

A considerable number of iodonium/diiodonium salts, mostly isolated from the final reaction mixtures in the form of sparingly soluble bromides/dibromides were synthesized, and their physical characterics, after the purification, as well as the IR and ¹H NMR spectra were reported. The following *tricyclic* aromatic systems were covered:

- 1) Dibenzofuran, 2- and 2,8-substituted with various aryliodonio groups [4].
- 2) N-Acyl-3-(Br, Cl or NO₂)carbazoles, 6-substituted with various aryliodonio groups [5].
- 3) Xanthene, xanth-9-one, fluorene and fluoren-9-one, substituted in their 2- and 2,7-positions with various aryliodonio groups [9]

Nearly all the iodonium salts synthesized were new. The aim of this work was to obtain several tens novel *iodine(III)* derivatives of the aforementioned tricyclic aromatic systems, which previously had scarcely ever been covered by the organic chemistry of polyvalent iodine. The *crude* iodonium bromides / dibromides obtained, after their effective oxidative metatheses into e.g. *pure* tetrafluoroborates (see Section 7.1), would next be reacted with various *nucleophiles* (inorganic and organic) to produce the corresponding mono- or disubstituted derivatives of the above tricyclic aromatic systems.

A large part of these novel iodonium salts were obtained with the short-cut method developed in our laboratory [3] (see Section 6.1). When *we failed* to obtain the desired iodonium salts using this method, we applied the other literature methods [72, 83, 84]; for more details see Refs. 4, 5 and 9.

6.3. Improved Syntheses of Some Diaryliodonium Salts from Symmetric Diarylmercurials and (Dichloroiodo)arenes (Willgerodt Method) [13]

Willgerodt [56, 86] had reacted cold (or hot [86a]) *aqueous* suspensions of *equal masses* [it practically means that the mercurials were used *in a deficit*] of powdered PhICl₂ with powdered Ar_2Hg (where Ar = phenyl, 2- and 4-tolyl, and 2-naphthyl) to afford the respective *diaryliodonium chlorides* (yields were not reported); sparingly soluble ArHgCl and other admixtures, e.g. PhIO [87], were hot-filtered off and discarded, viz.

$$\begin{array}{c} H_{2O} \\ PhICl_{2} + Ar_{2}Hg \xrightarrow{r.t., 12 h} Ph(Ar)I^{+}Cl^{-} + \bigvee ArHgCl \\ (excess) & (discarded) \end{array}$$
(Eq. 29).

Beringer and Lillien [88] applied the Willgerodt method to obtain three unsymmetric diaryliodonium chlorides. They obtained only *4-acetamidophenyl(phenyl)iodonium chloride* (which was precipitated out as its sparingly soluble *iodide*, isolated in **10%** crude yield) by reacting *equimolar amounts* of Ph₂Hg with 4-AcNHC₆H₄ICl₂ in hot water (40-50°C) for 12 hours. We obtained [13] the same *iodide*, but in **80%** crude yield, by reacting *equal masses* of PhICl₂ with symmetric 4,4'-mercuriobis(acetanilide) suspended in stirred hot water (40-50°C) for 12 hours. We explained this evident yield increase as follows:



Consequently, by reacting *equal masses* of PhICl₂ with symmetric 4,4'-mercuriobis(*N*,*N*-dimethylaniline), suspended in stirred hot water (40-50°C) for 12 hours, we isolated from the hot filtrate, after its cooling, *4-dimethylaminophenyl(phenyl)iodonium chloride* in **60%** crude yield. Previously, Beringer and Lillien [88] failed to obtain this iodonium salts, *para*-substituted with only one NMe₂ group; the same failure was also reported by Neiland [89]. A similar iodonium salt bearing the two *p*,*p*'-substituted NMe₂ groups was synthesized quite otherwise [90]; this synthesis is shown (Scheme 7) in our paper [13].

We also attempted, *without effect*, to synthesize various 8-(aryliodonio)caffeine halides with using the Willgerodt method. Hence, we used our short-cut, oxidative method [3] (see Section 6.1) to obtain 8-(4-methoxyphenyliodonio)caffeine bromide (49% crude yield) by acidic coupling of the oxidized 8-iodocaffeine with anisole. It is, in fact, the first *iodine(III)* derivative of caffeine, which may open novel routes for preparing 8-substituted caffeines by its reactions with various nucleophiles [21, 69, 84].

6.4. Short-cut Syntheses of Diaryliodonium Bromides with Sodium Perborate as Oxidant [96]

This novel method is just completed and will be published soon. Its essence is as follows:

(1)
$$\operatorname{ArI} + \operatorname{NaBO_3} \cdot \operatorname{H_2O} + 2\operatorname{Ac_2O} + 2\operatorname{H_2SO_4} \rightarrow \operatorname{ArI}(\operatorname{OSO_3H})_2 + [\operatorname{NaBO_2}] + 4\operatorname{AcOH}$$

- (2) $\operatorname{ArI}(\operatorname{OSO}_3\operatorname{H})_2$ (not isolated) + Ar'-H (in excess) \rightarrow Ar(Ar')I⁺HSO₄⁻ + H₂SO₄
- (3) Ar(Ar')I⁺HSO₄ (*not isolated*) + aq. KBr (*in excess*) $\rightarrow \downarrow$ Ar(Ar')I⁺Br⁻ + KHSO₄ (Eq. 30)

Powdered NaBO₃•H₂O (used in the 200% excess) was suspended in Ac₂O and this suspension was stirred for 30 min, at 30°C. Appropriate ArI (0% excess) followed by Ar'-H (used in the 400% excess) were slowly added at 40°C with stirring, and next the whole was cooled to 10°C. *Varied amounts* of conc. H₂SO₄ were slowly added dropwise with stirring while keeping the temperature below 15°C. The stirring was continued for a further 5-7 h while keeping the temperature at 15°C; some acidic couplings were carried out at 25-30°C. The mixtures were poured into ice water, followed by the extraction with Et₂O to remove the unreacted Ar'-H and other organic impurities. Excess aq. KBr was added to the aqueous layers to precipitate the respective bromides, Ar(Ar')I⁺Br⁻. They were collected by filtration, washed well on the filter with water, then with acetone and Et₂O, and air-dried in the dark; the *crude* bromides were obtained in 45-91% yields. Small crude samples were *quickly* recrystallized from boiling MeOH to give the analytically pure specimens.

7. Anion Metatheses in Diaryliodonium Halides

One of the most important applications of diaryliodonium salts is their use in the **arylation** of organic and inorganic nucleophiles [21, 69a, 84, 91], including the fluoride anion, which results in the

formation of *aromatic fluorides* [92]. It has been observed [21, 84, 91] that the best arylation yields of the nucleophilic bases are achieved by applying the diaryliodonium salts with substantially *non-nucleophilic* counterions, viz. tosylates, triflates, hydrogensulfates, perchlorates, etc.; the direct synthesis of diaryliodonium triflates has been published by Japanese workers [93]. Beringer and co-workers [91] have observed the superior yields with diphenyliodonium tetrafluoroborate in the phenylation of organic and inorganic bases; several diaryliodonium tetrafluoroborates were directly synthesized by Neiland [89]. The *sparingly soluble* perchlorates as well as *bromides, iodides*, picrates, tetraphenylborates, and 2,4,6-tribromobenzenesulfonates [72] were often nearly quantitatively precipitated out from hot solutions or suspensions of more soluble diaryliodonium salts with an appropriate counterion. However, a considerable number of necessary diaryliodonium salts were only attainable from the corresponding halides or hydrogensulfates by using costly and toxic lead(II), silver and barium salts, e.g. silver tetrafluoroborate, trifluoroacetate and nitrate, or lead(II), silver and barium organosulfates or organocarboxylates, or otherwise [21, 72, 84]. These approaches have been briefly reviewed [21b, 84].

Ionic inorganic and organic halides, except of fluorides, in hot aqueous or boiling alcoholic solutions (*acidified* with e.g. H_2SO_4) were more or less readily oxidized ($\Gamma > Br^- > CI^-$, fluorides do *not* react) by an excess of 30% aq. H_2O_2 , with evolution of the respective halogens [94]. We have applied this information to effect a variety of the **oxidative anion metatheses** in a large number of crude diaryliodonium *bromides* and *iodides*, which were formerly precipitated out from the final reaction mixtures with excess aq. KBr or KI solutions [3-5, 6b, 9, 13] (see Sections 6.1-6.3). This is explained below in Sections 7.1 and 7.2.

7.1. Oxidative Anion Metatheses in Crude Diaryliodonium Bromides [3]

In our paper [3] we have reported various **oxidative anion metatheses** in the *crude* diaryliodonium *bromides*, previously obtained by us with the one-pot protocols discussed in Section 6.1, which produced the corresponding *pure* diaryliodonium tetrafluoroborates, tosylates, trifluoroacetates, hydrogensulfates, nitrates, and chlorides in 57-80% yields. These new preparative procedures are easier and shorter, and also less expensive, than many earlier methods.

The essence of our oxidative anion metatheses is given below:

$$\overset{\bigoplus}{2 \text{ [Ar-I-Ar'] }} \overset{\bigoplus}{\text{Br}} + \text{H}_2\text{O}_2 (30\% \text{ aq.}) + 2 \text{ HX} \xrightarrow{15 \text{ min after complete dissolution}}{57-80\%}$$

$$\xrightarrow{\bigoplus} 2 \text{ [Ar-I-Ar'] } \overset{\bigoplus}{X} + 2 \text{ H}_2\text{O} + \text{Br}_2 (captured by cyclohexene)}$$
(Eq. 31).

 $X = BF_4$, TsO, CF₃COO, HSO₄, NO₃, Cl.

Eleven crude diaryliodonium bromides were suspended with stirring in pure MeOH (unless otherwise stated) mixed with the 100% excess of *cyclohexene* (acting there as a "halogen scavenger"), then an appropriate *strong* acid, HX (see above), was added with stirring, followed by 30% aq. H₂O₂ used in the 196% excess. The mixtures were stirred and boiled under a reflux condenser until complete dissolution of all the reaction components. The reflux was prolonged for a further 15 min. The solvent was distilled off under vacuum, and the crude products left in the flask were triturated with either Et₂O or acetone to remove any organic impurities. The following individual workups and recrystallizations resulted in the *analytically pure* iodonium salts, Ar(Ar')I⁺X⁻. For the metathesis into trifluoroacetates, a catalytic amount of ammonium molybdate was added to the reaction mixtures. For the metathesis into nitrates, 50% aq. AcOH at 70°C was used as the solvent, instead of boiling MeOH. For the metathesis into chlorides, a much slower competitive reaction also occurs [2HCl + H₂O₂ \rightarrow 2H₂O + Cl₂ (*captured by cyclohexene*)], hence somewhat increased amounts (by ca. 10%) of H₂O₂, HCl, and cyclohexene were used.

Note. Dr. Pawel Kazmierczak has recently established that *dry acetone* used as the solvent of choice in the oxidative anion metatheses of crude diaryliodonium *bromides* and *chlorides* (see Section 7.2) may act itself as a very efficient "halogen scavanger", hence the addition of *cyclohexene* to the respective reaction mixtures is not necessary.

7.2. Oxidative Anion Metatheses in Crude Diaryliodonium Iodides and Chlorides [6b]

The oxidative anion metatheses, similar to those reported in Section 7.1., in six *crude* diaryliodonium *iodides* and two *chlorides* produced the corresponding *pure* hydrogensulfates, nitrates, tetrafluoroborates, triflates, tosylates, as well as bromides and chlorides (only from the iodides) in 54-86% yields. By using the modified oxidative metatheses in the *iodides* (in the presence of HBr or HCl, but in the absence of cyclohexene) it was possible either to isolate, or to detect only, the intermediate *dihaloiodates(I)*, $[Ar_2I]^+[IX_2]^-$ (X = Br or Cl). A complex Ph₂I⁺Cl⁻•1/2I₂ was also obtained in 56% yield. Tetraethylammonium iodide, Et₄N⁺T, was similarly converted into pure tetrafluoroborate or dibromoiodate(I) in 75 and 76% yields, respectively.

The essence of our oxidative anion metatheses is as follows:

$$2 [Ar-I-Ar'] \stackrel{\bigoplus}{Y} + H_2O_2 (30\% \text{ aq.}) + 2 \text{ HX (a strong acid)} \longrightarrow$$

$$\xrightarrow{\text{ROH (reflux)}} 2 [Ar-I-Ar'] \stackrel{\bigoplus}{X} + 2 \text{ H}_2O + \text{Y}_2 \qquad (Eq. 32).$$

 $Y = I \text{ or } Cl; X = HSO_4, NO_3, BF_4, CF_3SO_3, TsO, as well as Cl, Br (only when Y = I).$

As previously Br_2 [3], also molecular chlorine, $Y_2 = Cl_2$, evolved in the oxidative reactions, was fully captured by the 100% excess of *cyclohexene* added on purpose as a "halogen scavanger"; 1,2dichlorocyclohexane as well as the other possible *cyclohexene adducts* with reactive agents (H₂O₂, Cl-O⁺H₂, etc.) were next removed from the crude metathesized products by their prior trituration with anhydrous Et₂O, followed by washing on the filter with the same solvent. For the diaryliodonium *iodides*, the diiodine, $Y_2 = I_2$, evolved was simply *washed off* from the crude metathesized products with anhydrous Et₂O. The following recrystallizations (see [6b] for details) gave *analytically pure* metathesized products.

Diaryliodonium *chlorides* were oxidized by H_2O_2 in acidified, boiling alcoholic solutions *much less readily* (3 h; 1076% excess of H_2O_2 added portionwise) than the respective bromides [3] (15 min; 196% excess of H_2O_2) and iodides (10 min; 36% excess of H_2O_2); the final yields of the respective *pure* metathesized products were 54-76, 57-80 [3], and 54-86%, respectively.

In our earlier work [3] we metathesized oxidatively two aryliodonium *bromides* into the respective *chlorides* with no comments. However, in our second work [6b] we found that such oxidative reactions are more complex than expected. This is likely to be due to a series of fast consecutive reactions taking place in the investigated solutions of diaryliodonium *iodides* containing H_2O_2 and acidified with excess HBr or HCl, but in the absence of cyclohexene, (a)-(d):

(a)
$$I^{\ominus}(only\ a\ part) + H_2O_2 \xrightarrow{H^{\oplus}} I-OH \xrightarrow{H^{\oplus}} I-OH_2$$

(b) $I^{\ominus}(unoxidized\ rest) + I-OH_2 \xrightarrow{-H_2O} I_2 (evolved; see Eq. 32)$
(c) $Br^{\ominus}(from\ excess\ HBr) + I-OH_2 \xrightarrow{-H_2O} IBr (transient) \longrightarrow \frac{Br^{\ominus}}{-H_2O} IBr_2^{\ominus}(intermediate)$
(d) $CI^{\ominus}(from\ excess\ HCl) + I-OH_2 \xrightarrow{-H_2O} ICl (transient) \longrightarrow \frac{CI^{\ominus}}{-H_2O} ICl_2^{\ominus}(intermediate)$

It is known that both diaryliodonium iodides and tetraalkylammonium iodides are easily converted into the respective dihaloiodates(I) either by the reaction of Cl_2 or Br_2 with the iodides, or by the reaction of IBr or ICl with the corresponding bromides or chlorides. For more details see Ref. 21a, Chap. 7.

By carrying out the oxidative anion metatheses in either diphenyliodonium iodide or tetraethylammonium iodide with excess HBr, we succeeded in the isolation of the intermediate, deep-orange colored, *dibromoiodates(I)* in 56 and 76% yields, respectively. The other diaryliodonium iodides gave only oily or semisolid, *deep-yellow* or *deep-orange mixtures* of dihaloiodates(I) with the respective diaryliodonium bromides or chlorides, when their oxidative metatheses were performed in the presence of excess HBr or HCl, but in the absence of cyclohexene. These mixtures may be easily converted into the unmixed bromides or chlorides in 86-92% crude yields by short (5 min) refluxing in anhydrous

MeOH admixtured with excess *cyclohexene* (see Eq. 33 below). This procedure for converting the dihaloiodates(I) into the corresponding diaryliodonium bromides or chlorides (X = Br or Cl) is more effective than that offered in the literature [91, 95]. This involved chlorination of the diaryliodonium iodides to the dichloroiodates(I) or tetrachloroiodates(III), which when dissolved in hot acetone reacted to give the iodonium chlorides.

$$\begin{bmatrix} \operatorname{Ar-I-Ar'} \end{bmatrix} \stackrel{\bigoplus}{I^{\ominus}} + 2 \operatorname{HX} + 2 \operatorname{H}_{2}\operatorname{O_{2}} \xrightarrow{20-30 \operatorname{min}} [\operatorname{Ar-I-Ar'}] \operatorname{IX}_{2}^{\ominus} \xrightarrow{} \operatorname{IX}_{2}^$$

where: HX = HBr or HCl; only *diphenyliodonium dibromoiodate(I)* was isolated in 56% yield as the intermediate. 1-X(Cl or Br)-2-iodocyclohexanes were *washed off* in full from the crude products, $Ar(Ar')I^{+}X^{-}$, with anhydrous diethyl ether.

Finally, we also performed some "model" study [6a] on quite similar oxidative anion metatheses in boiling methanolic solutions of KI, KBr, and KCl, acidified with H_2SO_4 , HBF_4 or *p*-toluenesulfonic acid. *Each* of the halides was effectively converted into pure potassium hydrogensulfate, tetrafluoroborate, and tosylate. This also confirms the usefulness of the presented method in *inorganic chemistry*.

8. Conclusions

This review shows our small research group's main interests in developing *novel* (or considerably improved) preparative procedures, which are suitable for easy, quick, cheap, effective, and possibly *environmentally benign* preparations of iodoarenes and some basic organic hypervalent iodine reagents: ArICl₂, ArI(OAc)₂, ArIO₂, and diaryliodonium salts with substantially *non-nucleophilic* counterions (suitable for the arylation of organic and inorganic nucleophiles). We hope that some of our methods will be applied as such in other organic laboratories and, possibly, in the pharmaceutical industry, but after their enlargements worked up in the experienced and well-equipped industrial laboratories; see e.g. Ref. 59.

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