After the solution had been kept at 0° for ten minutes, excess nitrous acid was removed by the addition of ammonium sulfamate. The volume at this point had increased to 25-30 ml. because of occasional additions of ice. N-(2-N-Methylanilinoethyl)-acetamide¹ (2.07 g., 0.0108 mole) was added and the mixture was stirred for one hour at 0-5°, after which 1.25 g. of sodium acetate trihydrate was added, and stirring was continued two hours. After the mixture had stood an additional two hours at 0-5° added, the mixture was stirred a few minutes, then left overnight in the refrigerator. After the slurry had stood at room temperature for one hour the precipitate was filtered off, washed with water and recrystallized from ethanol. The yield was 2.76 g. (86.5%) of orange-red parallelograms exhibiting occasional twinned forms with bifurcated ends: m. p. 177.5-178° (cor.). A solution of the substance in 0.1 N hydrochloric acid showed an absorption maximum in the visible spectrum at about 520 mµ.²

(2) Determination by Rachel N. Sclar, Cosmetic Division, U. S. Food and Drug Administration.

Anal. Calcd. for C₁₇H₂₀N₄O: N, 18.91. Found:³ N, 18.79.

Attempts to remove the acetyl group by refluxing with 6 N hydrochloric acid resulted in deep seated changes in the molecule,⁴ whereas heating with 5 N aqueous alcoholic alkali for seven hours gave only slight evidence of deacetylation.

In view of results repeatedly recorded for couplings involving N,N-disubstituted anilines having free p-positions, it is highly probable that the phenylazo group of this compound is in the p-position.

(3) By Thomas H. Harris, Food Division, U. S. Food and Drug Administration.

(4) Rowe and Dangerfield, J. Soc. Dyers Colourists, **52**, 48 (1936), and Hodgson and Foster, J. Chem. Soc., 755 (1941), have reported on the decomposition of certain azo compounds by mineral acids.

CHEMICAL SECTION, DRUG DIVISION

U. S. FOOD AND DRUG ADMINISTRATION

FEDERAL SECURITY AGENCY LLEWELLYN H. WELSH WASHINGTON 25, D. C.

RECEIVED AUGUST 31, 1945

COMMUNICATIONS TO THE EDITOR

Sir:

DIETHYLSILANEDIOL

Although organosilanediols have been known for forty years^{1,2} no dialkylsilanediol has been reported. The non-isolation of this type compound was generally attributed to rapidity of intermolecular condensation.³ Hydrolysis studies on diethyldichlorosilane have previously failed to give the diol.⁴

By careful control of conditions during the hydrolysis and the working up of the product we have prepared diethylsilanediol in good yield from diethyldichlorosilane. Best results were obtained with dilute alkali or saturated sodium bicarbonate solution. However, concentrated sodium chloride solution may also be used. A solution of 30 g. of diethyldichlorosilane in 500 cc. of dry ether was added during five minutes with stirring to the theoretical amount of 1.5 N alkali at 0° Immediate separation of the ether layer and ether extraction of the water layer were followed by drying over 25 g. of potassium carbonate for a short time. The ether solution of the product was then concentrated in vacuo to 250 cc. Addition of 250 cc. of pentane followed by cooling gave 15 g., 65% yield, of diethylsilanediol, a white

(1) For references to diphenylsilanediol see Burkhard, THIS JOURNAL, 67, 2173 (1945).

(2) For the preparation of the phenylethyl, benzylethyl, benzylphenyl and dibenxyl compounds see Robison and Kipping, J. Chem. Soc., 101, 2148 (1912); Robison and Kipping, *ibid.*, 101, 2156 (1912).

 (3) See for example Krause and Grosse, "Die Chemie der Metallorganischem Verbindungen," Borntraeger Geb., Berlin, 1937, p. 284; Rochow, Chem. Eng. News, 23, 614 (1945).

(4) Martin and Kipping, J. Chem. Soc., 95, 313 (1909); Hyde and DeLong, THIS JOURNAL, 63, 1194 (1941).

crystalline solid, m. p. 96° before and after recrystallization and boiling with decomposition at about 140°.

Anal. Calcd. for C₄H₁₂SiO₂: Si, 23.34; C, 39.97; H, 10.07. Found: Si, 23.12; C, 39.91; H, 10.10.

Diethylsilanediol can be stored in ordinary corked bottles. Even exposure to a laboratory atmosphere for long periods of time gives no change. At room temperature it is soluble in water and in oxygen-containing organic solvents, and is sparingly soluble in chlorinated hydrocarbons. The compound is easily recrystallized from ether-pentane, acetone-pentane, chloroform and carbon tetrachloride.

Treatment of diethylsilanediol with thionyl chloride and quinoline gave diethyldichlorosilane. Refluxing with benzene and iodine gave the theoretical amount of water. More extensive data on the preparation and properties of dialkylsilanediols will be reported shortly.

We thank R. N. Walter of this Laboratory for the analyses reported in this Communication.

DEPARTMENT OF CHEMISTRY THE PENNSYLVANIA STATE COLLEGE STATE COLLEGE, PENNA. RECEIVED JANUARY 10, 1946 P. A. DIGIORGIO L. H. SOMMER F. C. WHITMORE

o-NITROBENZOYL CHLORIDE AND HAZARD INVOLVED IN ITS DISTILLATION

Sir:

N. C. Cook and F. C. Whitmore have called attention (*Ind. Eng. Chem.*, *News Ed.*, 23, 2394 (1945)) to the treacherous distillation behavior

⁽¹⁾ J. von Braun, et al., Ber., 70B, 979 (1937).

of o-nitrobenzoyl chloride. Our experience was similar. Small quantities appeared to distil satisfactorily, but this trouble was encountered in an 8-mole run. The volatile products (POCl₃, PCl₅) were removed at 100° (20 mm.), then the flask was equipped for vacuum distillation with an oil pump and a glas-col heating mantle. Before any distillate was collected the material detonated violently. Fortunately, however, there was a warning period of a few seconds.

Reasonably pure *o*-nitrobenzoyl chloride may be made without distillation by use of thionyl chloride instead of phosphorus pentachloride. The gaseous reaction products and any unused thionyl chloride were removed by aspirating with a stream of dry illuminating gas for a few hours.

DEPARTMENT OF CHEMISTRY Northwestern University Evanston, Julinois	W. A. BONNER CHARLES D. HURT
LYANSION, ILLINOIS	CHARDES D. HORL
RECEIVED JANUARY 9,	1946

CASHEW NUT SHELL LIQUID. II. AN APPLICA-TION OF THE PREVOST REAGENT TO ALKENYL PHENOLS¹

Sir:

The noxious saps of numerous members of the *anacardiaceae* family, such as Poison Ivy, Cashew Nut Tree, Japanese Lac Tree, *etc.*, contain phenolic mixtures made up of saturated, mono-, diand probably polyolefinic components. With the purpose of developing a method for separating and establishing the structures of such olefinic components, we have investigated the use of silver iodobenzoate.^{2,3}

The monophenol, obtained by direct vacuum distillation of raw commercial cashew nut shell liquid, and possessing an average of two aliphatic double bonds,¹ was redistilled several times, methylated and then distilled. The resulting methyl ether possessed 1.56 double bonds. Treatment with silver iodobenzoate, and subsequent hydrolysis of the benzoates, yielded, on fractional crystallization from aqueous methanol, two crystalline glycols (A and B), both of which contained iodine. Both glycols showed no discoloration of bromine in carbon tetrachloride.

Glycol (A), m. p. 92–93°, analyzed correctly for an iodinated monoglycol.



Anal. Calcd. for $C_{22}H_{37}O_3I$: C, 55.35; H, 7.74; mol. wt., 476. Found: C, 55.55; H, 7.51; mol. wt. (ebulliometric in CCl₄), 478.

Glycol (B) m. p. 112–113° and analyzed correctly for an iodinated diglycol.

Anal. Calcd. for C₂₂H₃₇O₅I: C, 51.96; H, 7.28. Found: C, 51.92, 52.01; H,7.50, 7.43.

Oxidation of the mono glycol (A) with periodic acid in aqueous methanol gave *n*-heptaldehyde which was identified as the 2,4-dinitrophenylhydrazone, m. p. $106.5-107^{\circ}$, by mixed m. p. and analysis.

Anal. Calcd. for C₁₃H₁₅O₄N₄: C, 53.06; H, 6.12; N, 19.05. Found: C, 53.23; H, 6.21; N, 19.10.

Oxidation of the mono glycol (A) with alkaline aqueous permanganate gave an aromatic acid containing iodine which analyzed correctly for (C).

Anal. Calcd. for C₈H₇O₈I: C, 34.54; H, 2.51. Found: C, 34.71; H, 2.77.

Catalytic hydrogenation of the mono glycol (A) using 5% Pd-BaCO₃ in ethanol yielded an iodine-free solid which is now under investigation.



The experiments described above establish that the mono phenol of commercial cashew nut shell liquid is a mixture of at least two different olefinic components: one a mono-olefin and the other a di-olefin. It appears likely that a more highly unsaturated component is also present. The monoolefinic component is unsaturated in the 8–9 position of the side chain as established by the isolation of η -heptaldehyde after periodic acid oxidation of the glycol.

Further studies on the identification of the position of the iodine in the ring and the application of the Prevost reagent to the separation and identification of other unsaturated natural phenolic bodies, such as are found in poison ivy, are now in progress. Details of all this work will appear in forthcoming publications.

DEPARTMENT OF CHEMISTRY COLUMBIA UNIVERSITY New York, N. Y. CH

IVERSITY MEYER SLETZINGER J. Y. CHARLES R. DAWSON RECEIVED JANUARY 16, 1946

SELECTIVE DEMETHYLATION OF PARAFFIN HYDROCARBONS

Sir:

Paraffin hydrocarbons, as for example, pentane and hexane, are known to react in the presence of active hydrogenation catalysts, such as nickel, to produce primarily methane and carbon. Mixtures of hexane and hydrogen give methane as the major product at 400°.

We wish to record a new reaction, the selective demethylation of branched-chain paraffin hydro-

⁽¹⁾ For the first article in this series, see Wasserman and Dawson, Ind. Eng. Chem., 37, 396 (1945).

⁽²⁾ Prevost, Comp. rend., 196, 1129 (1933); 197, 1661 (1933); 198, 2264 (1934).

⁽³⁾ Hershberg, Helv. Chim. Acta., 17, 351 (1934).