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## NEUROLDGICAL

The first of two articles written for Res Medica by J. H. STANTON, F.R.C.P.E., F.R.C.P., D.P.M., Neurological Unit, Northern General Hospital.

Wilfred Trotter has said that the performance of a refined neurological examination is "a job for men". Certainly performing a full neurological examination seems to separate the men from the boys and many medical students are often unnecessarily alarmed at the prospect. Much of this anxiety can be dispelled, however, if the logic of the examination is approciated. A greater number of objective signs can be clicited in the examination of the nervous system than in any other system and this profusion of signs, at first so umnerving, can be a positive advantage in providing sufficiently precise information regarding the site of dysfunction in the nervous system. After the examination has revealed this anatomical diagnosis, the physician, by taking into account the details of the cololution of the discase revealed in the history, can usually reach a final conclusion regarding the nature of the disease which is causing the dysfunction in the nerrous system. This is the final or pathological diagnosis.

The following account deals briefly with the
correct methods of eliciting the physical signs and is not primarily concerned with neurological history taking. The importance of taking a careful history, however, cannot be over-mphasised, in neurology as in any other branch of medicine. A well-known neurologist has said that if he does not have a pretty shrewd idea of what is wrong with a patient when he puts his pen down at the end of taking the history he probably never will know. Not only docs the history reveal important information, but the intelligent use of this information will often direct the neurologist's attention to those parts of the physical examination which are particularly revealing in the patient concenned.

## EXAMINATION OF THE CRANIAL NERVES

First cramial nerve. The sense of smell should be tested by the use of various aromatic odours, such as coffee, oil of almonds or peppermint. The method is to occlude one nostril
and to ask the patient to sniff the sample presunted through the other nostril. This procedure is repeated on the opposite side. The patient may smell nothing (anosmia), or he may say that he can suchl some odour but is mable to identify it (hyposmia), or he may b. able to ickentify the odour correctly, in which case he has certainly got normal olfactory sense. 'lhe sense of small may be lost unilaterally with lesions of the olfactory tract and is not uncommonly lost altogether following haded injury.

The second cramial nerve. There are thate aspects of the function of the second cranial werve which have to be tested. I'hese are the vistal acuity, the ficlds of vision and the appearance of the funclus. Visual acuity is tested ideally for both far and near vision. At the bedside. however, usually only near dision is tested. To do this the patient covers onc cye with his hand and is given the nearvision type card and is requested to read the smallest type that he can sec. This is $\mathrm{N}_{5}$ on the standard near-vision test. Each eyc is tested separately. The ficlds of vision are tested by confrontation. The examiner faces the patient at a distance of $18^{\prime \prime}$ to $2^{\prime}$. One of the patient's eyes is covered and he is instructed to look at the opposite eve of the examinet. In this way the patient will be looking with, for cxample. his left eye into the examiner's right eye. The pationt is instructed to keep his eve fixed on the examiner's eye and the examiner then proceeds to test the extent of the patient's peripheral field by direct comparison with that of his own cyc. A small object such as the head of a hatpin is introduced from the periphery towards the centre matil the patient declares that he can see it. 'This precedure is carried out around the whole circumference of the field of vision and any constriction of the field or sector defect is readily apparent. The method of confrontation can also be used to determine the presconce of a central scotoma. In this case the object is held a few degrees outside the line of fixation of the patient's ere and he is asked to say whether the object appears clearer in this position or when it is brought immediately in front of his fixation point. Nomally the latter position is, of course, the clearer. The cxmmation of the fundus repaires the use of the ophthalmoscope and this can only be acquired by considerable practice. The examination of the fundus is. of course, part of the general physical examination, but from the
ncurological standpoint certain fealures are cspecially important. These are (a) the appearmee of the dise: 'This is normally of a pinkishwhite colour, paler than that of the surromeling retina, and in the centre of the dise the optic cup can be seen. from which the retinal ressels emerge and rum over the dise to reace, the periphery of the retina. In optic alrophy the clise is paler than normal, and the eelge sharper, and in rased intracranial pressite. papillocedema occurs in which there is mative swelling of the optic disc. Suelling can be suspected when the dise is pinker than nomal, and when the optic cup is filled up and when the edges of the clise become blared as the ocdema masks the junction with the sumomeding retina. (b) The vessels of the retum: Cancful examination may reveal diffecomes in calibre of the arteries, and in the artemo venems ratio, indicating the presence of ations selerosis, or may show the presence of anterial or venous thrombosis. (c) The appearance of the retina itself: 'The presence of hamontages or exudates is of importance, since these ocen not only in general medical discases. sued as hypertension, renal discase, and diabets, but will also oceur when there is gross rained intractanial pressure with papillocdema. The retina may also show sigus of old choroitlitis or other infammatory discase, such as tuber. culoma. toxoplasmosis, cte., which may bave a bearing on the newrological condition.

Third, fourth and sixth cramial nerees. di this point it is comenient to cxamine the on bit for the presence or absence of proptosis, as well as testing the integrity of the musele; mnervated by the third, fourth and sixtio canial nerves, which also include the levator of the upper lid and the pupil. Proptosis is last detected by examining the patient from behind. Ptosis is usually noted carly on meceng the pationt and is comeniently moasmed be the degree to which the upper lid in the :omil open-eyed position covers the upper part of the limbus. 'To examine the ocular morements the patient is first asked to look directly athead. In this position it is possible to cletect any squint, that is to say to eletect whether the visual axes appear paralled or not. The pateme should then be asked to fix his gaze on the examiners finger licld at a distance of alout $z$ ' from the patient. 'Ihe finger is then :noveci with the patients cres following it to the extreme positions of lateral gaze to cither side and also vertically upwards and clownards. The patient is asked whether he sees dunble
in any position of the eres and if he does the direction of gaze which produces mamaman separation of the images is determined. it 1 ?: extremes of gaze, attention should also be paicl to the presence of nestagmus.

The pupils: These are normally equal in size and regular in shape. 'They should constrict to a light stimulus, dircet or consensual. and on accommodation/consergence. 'The light reflex is tested by covering one eye and shiming a lighted torch on the other ere. 'Th: pupil of the eye so stimulated should contrate and remain constricted so long as a lighe is shone on it. If at the same time the hand covering the other ere is raised, it will be seen the pupil of this cye nomally contracts consensually when liglit is thrown on the otler cye. 'lhis procedure is carricd out on cach side. Accommodation/ comergance is tested be asking the patient first to look into the distance and then to look at the observer's finger placed one foot in front of him. The two exes should converge and the pupils contract when the patient focuses on the finger.

Fifth crmial nerec. The sensory, motor and refles functions of this nerve must be tested. Scusation to light touch and pin prick is examined in the territory of the thene divisions of the nerice. Motor function of the muscles of mastication, the masseters, temporales and pterygoids, is also tested. 'l'his is done by asking the patient to open or close the jaw against resistance. If the muscles of mastication are paralysed on one side, the jaw will deviate towards that side when opening. 'The reflexes of the trigeminal nerve are the corncal reflex, which is tested by lightly applying it wisp of cotton wool to the corica, and the jaw jerk which is the tendon refles of the muscles of mastication. The patient is asked to open his jaw half-way and the jaw is grasped between the thomb and forefinger of the cxamincers hand. A blow with the patella hammer on the cxaminer's thumb will then clicit the jaw jerk.

The serenth cramial nerve immervates the muscles of the face. It also carries the sensation of taste from the anterior two-thirds of the longuc through the chorda tympani nerve. The facial museles are tested by asking the patient to wrinkle his forchead, to frown, to serew his eyes up tight, to smile and to whistle. In upper motor nemone weakness the morements of the lower part of the face only are
involved, but in a lower motor neurone palsy the whole of the musculature on the affected side is paralysed. I'aste is a relatively enude sensation and comprises the distinction of sweet, sour, salt and bitter. Other more delicate flavours which we loosely call taste are in fact the propertics of the first cranial nerse. Taste on the tongue is tested by applying a small amount of sugar, citric acid. salt or guinine with an applicator stick to the protruded tonguc, the patient being asked to identify the taste without withdrawing his tongue into his mouth. I Ie should be instructed to raise his finger to indicate when he has appreciated the taste. I le can then be allowed to withelraw his tongue and to anomece what he tasted.

Eighth cramial norec. llearing is lested it the bedside by the distance at which the patient can hear the whispered voice while one ear is ocelucled. Normally the distance should be over three feet. If there is depression of hearing of one or both cars then the tuming fork tests should be carried out. 'lhere are two of these tests: 1) Rimecs test consists of a comparison of bone and air conduction of sound. The vibrating tuming fork is placed against the mastoid process and the pationl asked to say as soon as he can no longer hear the note. 'The tuming fork is then remored and the vibrating end presented close to the cxternal anditory meatus when, if air conduction is (as nomally) better than bone conduction, the patient will again hear the note. This procedure is carried out on cach side. (2) Weler's test is performed by placing the vilorating toming fork on the midline of the head either at the forehcad or the vertex and asking the patient whether he hears the sound. With normal hearing the sound is described as in the middle of the head or all over the head, but when there is impaired hearing the sound may be localised to one side. If the deafness is associated with depressed air conduction and Weber's test is lateralised to the same side, this indicates a middle-car deafness. If in the deaf ear air conduction is better than bone conduction. while both are elepressed. and Weber's test is lateralised to the nomal ear, then the deafness is caused by nerve deafucss.

The winth and tenth cramial norves innervatic the soft palate and the muscles of the pharyme and largn. 'l'le minth nerve also carrics the
sensation of taste for the posterior third of the tonguc. The function of these two nerves can be tested by obscrsing the movements of the palate when the patient says 'Ah', and by cliciting palatal and pharyngeal reflexes. Touching the soft palate with an applicator stick causes a brisk contraction with elevation of the palate, and touching the posterior pharyngeal wall causes gagging with contraction of the wall of the pharynx. If there is paralysis of one side of the palate or pharynx then these movements will be assymmetrical, the movement appearing only on the normal side. Taste over the posterior third of the tongue is tested as described under the facial nerve.

The eleventh cramial nerve imnervates the sterno-mastoid and the upper part of the trapezius muscles. The sterno-mastoid is tested by asking the patient to turn his head to the opposite side against resistance when the
contracting muscle can be seen and its power cstimated. The trapezius is tested by asking the patient to shrug his shoulders upwards towards his ears, while the examiner presses down on the shoulders to assess the power.

The twelfth cramial nerve, the hypoglossal, innervates the tongue. The tongue should first be inspected for any muscle atrophy, fibrillation or tremor. The patient should then be asked to protrude his tongue which normally protrucles in the miclinc. If there is paralysis of one side then the tongue will deviate towards that side. If this paralysis is of upper motor neurone type, there will be no wasting of the tonguc. It it is of lower motor neuronc origin, there will be atrophy and possibly fibrillation of the paralysed side.
(To be continued)

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# THE FIELD OF ALLERGY 

by K. K. Adjepon-Yamoah, B.Sc.

## A review based on a dissertation read before the Society on 10th December, 1965.

'This article concentrates on (i) the immunological basis, (ii) pathophysiological mechanisms, and (iii) control (theoretical and practical) of the immediate-type allergy.

## PART I - IMMUNOLOGICAL BASIS

## Introduction

Antigen-antibody reactions constitute an important group of defences, facilitating phagocytosis and blocking the toxic effects of parasitic poisons. The reaction confers 'immmity'. 'The combination of antigen and antibody is, however, not always beneficial. Pathological reactions as severe or more severe than the cffect of the antigen alone are sometimes noticed. Hypersensitivity or allergic reactions form major cxamples of such conditions. Allergy or hypersensitivity may be defined as a state in which the animal reacts in an excessive way to the introduction of an antigen or a hapten even though the antigen or hapten may be imnocuous. Not all instances of hypersensitivity enjoy the ielentification of the exciting antigens, the mediating antibodics and the mechanisms of tissue damage.

## Classification

Experimentally two types of hypersensitioity reactions can be demonstrated. They are:
(a) the 'immediate type' and (b) the 'delayed type' reactions. Some of the main differences are summarised below.

|  | Immediate Type | Delayed Type |
| :---: | :---: | :---: |
| 1. Speed of onset of reaction following antigen introduction | Immediate | Delayed 24-72 hours |
| 2. Type of antibodies | $\gamma$-globulins | As yet Unidentified |
| 3. Chemical mediators | Histamine 5 HT, SRS-A, ? Bradykinin. (Depending on species) | As yet Unidentified ? Bradykinin |
| 4. Transfer from animal to animal | Possible with serum in many instances | Not possible with serum. Possible with cells in animals. In man extracts of cells are effective. |
| 5. Types | Anaphylactic shock, Arthus reaction serum sickness, atrophy e.g. asthma, some drug sensitivity, Allergic rhinitis. | Bacterial allergy e.g. Tuberculin type reaction. Contact sensitivity to simple chemical, e.g. contact dermatitis, |

## Anaphylaxis

Portier and Richet (1902) found that whereas the first intravenous injection into dogs of an extract of sea ancmones was relatively harmless, a second injection some 2 weeks later resulted in violent symptoms and often in the death of the dogs. Instead of 'phylaxis' (i.c. immunity), anaphylaxis developed. Soon Theobald, Smith, and Otto independently slowed that the guinca pig could likewise be made hyperscisitive even to non-poisonous extracts. To explain these facts two schools of thought developed. The first and now defunci theory was the 'anaphylatoxin hypothesis' led by Portier and Richet. The second and now widely accepted theory is the 'cellular hypothesis' supported by Dale. Bricfly, this cellular theory maintained that the anaphylactic reaction was the result of union between antigen and antibody which had become 'fixed' to the living cell surface - Dale and Schultz indcpendently showed that the phenomenon of anaphylaxis could be demonstrated in isolated tissues without the presence of blood. Dalc showed that the uterus of a sensitised guinca pig (i.c. guinea pig which had reccired smiall injections of egg ovalbumin three weeks carlicr), when suspended in a nutrient fluid at $37^{\circ} \mathrm{C}$ and oxygenated, would contract upon the addition to the bath of a small amount of the substance against which the guinca pig had been sensitised. The effect was quite specific since unrelated antigens gave no reaction. Furthermore, after the uterus had once responded by contraction to the antigen in question, a second addition of the same amount of the same antigen produced no effect. The tissuc had thus become descusitised. Other workers have shown that other smooth muscle strips from sensitised guinca pigs behave in the same 'Schultz-Dalc' manner as the myometrium. It is now possible to scnsitise guinca pig tissucs passively by soaking them in antibody solution (e.g. $\mathrm{I}^{133}$ labelled cgg ovalbumin). There is ample evidence that antibody fixation to certain tissucs is a necessary prercquisite for anaphylaxis.

## Dcsensitisation

Guinca pigs sconsitised to maphylactic shock can be descensitised by repeated small injections of antigen. It las been shown that during the process of desensitisation a very high titre of circulating antibody is produced, and if antigen
is administered to such an immune animal, amaphylactic reaction docs not develop because the anount of circulating antibody is sufficient to neutralise all the injected antigen. Serum from such an immune animal is capable of inducing 'passive' scmsitisation to anaphylaxis, thus demonstrating that the antibodies involved are of the same type.

## Specics variation

There is considerable species variation in the manifestations of gencralised anaphylaxis. In the guinca pig there is severe bronchospasm leading to asphysia and death. In the rabbit death is ascribed to corpulmonale. In the dog, death is duc to hepatic congestion and peripheral circulatory failure. Man resembles the guinca pig in that there is acute respiratory distress of asthmatic type and generalised ocdema. Anaphylactic reaction in man is in fact rare but when it docs occur, it usually follows repeated injections of therapeutic serum (c.g. ATS), or cortain drugs (e.g. peniicillin and ncoarsphenaminc).

There is experimental cyidence that cot cleaths, which are responsible for about 2,000 infant deaths per year, may be due to hyperscnsitivity to cow's milk protein.
The disturbances in generalised anaphylaxis are fundamentally the same in all species. The main effects are: (a) spasm of smooth muscle, and (b) damage to endothclium of blood ressels and an increase in permeability, giving rise to gencralised ocdema.
Anaphylaxis has been used as an experimental model in the study of allergy. The basic mechanisms are not fundamentally different from other types of immediate hypersensitivity.

## Atopy, Food, Dust and Drug Scnsitivity

This group of allergics occur after the ingestion of certain foods and drugs, the inhalation of antigens like pollen, and the injection of drugs. There is considerable variation in symptomatology which seems to be dependent upon the route of absorption and the nature of the antigen or hapten.
Examples:
(a) Inhalation antigens, e.g. pollen, gives rise to respiratory symptoms such as allcrgic rhinitis, hay fuer and asthma.
(b) Ingested substances, e.g. mushrooms, shellfish, give rise to gastro-intestinal symptoms and rashes. There is possibly absorption of
whole protein from the gut, so providing an antigen.
(c) Injected drugs, e.g. streptomycin and penicillin, usually give rise to skin rashes.

There seems to be a genctic basis in these types of allergy - hence the mame atopy.

## Miscellancous Examples

Other examples of immediate hypersensitivity are Arthus reaction and serum sickness, but these conditions seem to be dependent on antigen-antibody complexes.

Many discases have been labelled allergic although their pathogencses are by no means clear. Examples are Type I ncphritis, rheumatic fever and polyarteritis nodosa. A number of drug 'diseases' have also been documented as being allergic, and chlorpromazine obstructive jaundice is a well known example of this group.

## PART II - MECHANISMS


'The mechanism by which antigen-antibody combination brings about the relcase of pharmacological agents is far from being well understood. Briefly - antigen combincs with fised antibody. This 'reaction' is believed to lead to activation of tissuc enzyme systems which include chymotrypsin-like enzymes and phospholipasc A (Austen and Brocklchurst. ig(u, etc.). Complement may or may not play a part at this stage.

Activated enzyme systems cause changes in the cells, such as mast cells which relcase pharmacologically active substances notably histamine, heparin, SRS-A, ; HT, and bradykinin. The pattern of release is to some extent elcpendent on the specics. The activation of tissue proteases and esterases may act on substrates such as pepticles in the blood to relcase rasoactive substances such as kinins.

The symptoms of hypersensitivity result from the actions of these pharmacological agents. A summary of the cvidence supporting the above statements is made below.

## Ensymic Participation

The influence here stems from indirect cridence in which the effects were observed of pH change, temperature change, calcium lack and specific enzyme inhibitors on certain standard tests, c.g. Schultz-Dale type of test. Mongar and Schild concluded (1062) that the cnzymes werc calcium reguiting and heat labile.

## Role of Histamine

As Schachter states "Ever since that time that the similarity between the symptoms of histamine intoxication and acute anaphylactic shock was pointed out by Dalc and Laidlaw (1910) an impressive body of cvidence implicating histamine in anaphylaxis has accumulated." Many workers have demonstrated the release of histamine from sensitised organs both in vitro and in situ by specific antigen. Histamine liberators, e.g. $48 / 80$, are able to reproduce many of the symptoms of anaphylactic shock when administered to animals. Schayer, and others, using radioactive histidine, have concluded that not only do mast cells store histaminc, but also form histamine from histidine. Extrision of mast cell granules, which are thought to contain histaminc-heparin complexes, have been observed during antigenantibody reaction. The evidence for the release of histamine in anaphylaxis is overwhelming and the relcase of this substance has been assumed to occur also in other immediatetype allergic reactions.

## Slow Reacting Substances of Anaphylaxis (SRS- - )

Kellaway and Trethewic (1940), reported the occurrence of a slow reacting substance from a sensitised tissuc following a challenge
with an antigen. The perfusate from guinea pig lung was assayed on guinca pig ileum and these workers recognised that the contraction differed from that caused by histamine in that the gut was slower to rclax.

Brocklchurst (1952) noticed that high concentrations of antihistamines were unable to abolish SRS-A responsc. SRS-A does not appear to exist in preformed state, but is gencrated by events set in motion by antigenantibody 'rcaction' (Brocklehurst). In the tissucs of sensitised guinca pigs and in human asthmatic lungs challenged with the appropriate antigens SRS-A is relcased along with histamine, but the peak relcase of SRS-A occurs later than that of histaminc and moreover the relcase of SRS-A continucs longer. SRS-A can cause a strong and well-maintained contraction in isolated human bronchioles and it is presumed to play an important rolc in asthma and so to be as least part of the cause of therapentic failures of antihistamines in this condition. Hersheimer and Stressman (1961) have shown that whereas impure SRS- $\Lambda$ acrosol decreased the vital capacity in asthmatic patients, it had only a small effect in normal subjects.

## Other Substances

5 HT has been shown to be important in some species (rabbit and mousc) but not in man.

Bradykinin is present in the blood cluring anaphylaxis in several species of animals and can mimic some of the changes which are not abolished by antihistamines and presumably cannot be attributed to histaminc. An enzyme capable of forming braclykinin in plasma, from dog plasma pseudoglobulin and from Mawer Fraction C is rapidly relcased from sensitised guinca pig lung or skin when these blood-frec tissues are challenged with specific antigens (Brocklchurst).

Some of the inflammatory changes accompanying antigen-antibody 'interaction' might be duc to bradykinin generated locally.

## PART III - CONTROL OF ALLERGY

Theoretically the allergic reaction can be prevented in a mumber of ways:-
(a) The first anti-allergic step, often impractical, is the aroidance of contact with known antigens.
(b) By preventing antibody synthesis, c.g. by total body irradiation, antimetabolites and corticosteroids. The obvious disadrantages here far out-weigh any possible therapeutic advantages.
(c) By preventing antibody fixation to tissucs. (This has not been possible.)
(d) By inhibiting enzymes involved in the allergic process. Little is known about these ccllular cozymes, although it is possible they have normal physiological functions, and so it follows that inhibition of these enzymes may interfere with some vital metabolic processes (Brocklchurst, 1962).
(c) By desensitisation. This method has been tried, but the results are often disappointing even when the existing antigen has been indentified.
(f) The last, and at present most simple method of controlling the allergic symptoms is the imhibition or elestruction of the pharmacological substances relcased during antigenantibody 'reaction'.

There is no satisfactory way of antagonising SRS-A although it has been reported that homochlorcyclizine is a uscful therapcutic agent in sceveral allergic conditions including asthma. This drug has multiple actions anti IITT, antihistaminc, antiacctylcholine and weakly anti-SRS-A. It is therefore difficult to predict which action is responsible for the clinical improvement.

Theoretically there are three ways in which a drug can oppose the actions of histamine:
(i) by plysiological antagonism - c.g. adrenaline which has many of its phamacological actions opposite to those of histamine.
(ii) the drug might elestroy histamine, e.g. fomaldehyde, nitrates and the enzyme diamine oxidase. These drugs are of very limited therapentic value.
(iii) by preventing histamine from reaching its site of action, e.g. by competition - the antihistamines. The last measure has proved to be the best therapeutic method of controlling the allergic reaction at present.

## Antihistamines

These drugs oppose all the effects of injected histamine exeept that on gastric secretion. The use of antihistamines in allergy, however, has certain serious disaduantages. I shall summarise these under three short paragraphs.

Disadvantages of the drugs. These include multiple actions of these antihistamines. All of them depress and sometimes stimulate the C.N.S. About $20 \%$ of people on antihistamines complain of minor and moderately severe side effects. Few complain of serious side effects such as blood dyscrasias, the paradoxical oceurrence of hypersensitivity reactions.

Therapeutic uses of antihistamines. The best therapeutic results have been obtained in acute uticaria and seasonal hay fever. In perennial vasomotor rhinitis, chronic urticaria, angioneurotic ocdema and allergic reactions to various allergens including drugs, the results of antilistamincs are less gratifying. In serum sickness they are of symptomatic value.

For acute anaphylactic reactions the antihistamines are not as effective as adrenaline or the corticosteroids. Antihistamines have failed to bencfit patients with bronchial asthma in spite of the undoubted allergic basis of the condition. Again these drugs have not been shown to be of any therapentic valuc in the so-called allergic discascs like polyarteritis nodosa, acute rheumatic fever and type I nephritis.

Possible interference with a physiological role of histamine. Histamine is widely distributed in the organism. It is stored in a readily relcasable form. Again, Schayer has shown recently that histamine is reaclily formed ceren from tissuc free of mast cells in response to stress. He has postulated a microcirculation regulator role for 'inducable' histamine (which is perhaps stretching conclusions too far). Kahlson (1962) has also shown increased production of 'nascent' histamine in rapidly growing tissucs, c.g. granulation tissuc, tumour cells and rat embryos. Many other obsersations have forced Kahlson to conclude that
'nascent' histamine formation is an intcgral part of the metabolism of certain rapiclly growing tissucs. I find it hard to believe that the presence of histamine in the complicated and homeostatically balanced organism is simply to cause pathological changes. A physiological role for histaminc is a rery distinct possibility and has to be studied. Blocking the actions of histamine in certain clinical situations may have other dangers and possibly what one gains on the romndabout is lost on the swings.

## PART IV - CONCLUDING REMARKS

The immunological basis of the allergic process is undoubtedly proved, but the reaction sequence after antigen-antibody combination is. still not clear. Progress has been made in the study of the pharmacological agents thought to be responsible for the allergic symptoms, but here too, there are gaps. Why should the protective action of antigen and antibody combination result in detrimental reactions in certain individuals? Is this duc to an imborn crror? If so, what is the basic biochemical abnormality that is involved? What is the agent responsible for the propagation of the allergic process once started? The fertile soil of the delayed-type hypersensitivity has not yet been cultivated.

It is cstimated that about one person in ten in Great Britain suffers from one kind of allergy. It is also thought that in spite of individual susceptibility anyone is liable to develop a type of hypersensitivity reaction if exposed to the antigen for a certain time. Allergic ractions to drugs and antitoxins pose more problems in therapeutics. Many of these considerations given above make a thorough understanding of the allergic process highly desirable.

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## RES

# MEDICA 

ABORTIVE LEGISLATION?

For the back-street abortionist business is booming. It is commonly estimated that some 100,000 criminal abortions are performed annually in England and Wales alonc. Maternal deaths from these are, according to Goodhart (1964) of the order of 35 per year in England and Wales, gising a surprisingly low maternal mortality of 0.35 per 1,000 which equals maternal mortality from all other causes. Serious maternal morbidity, however, defies estimate but must be alarmingly high. Eren in hospital, the operation carrics serious risk of complication, which is obviously greatly increased in a tenement kitchen. Cervical incompetence, causing repeated miscarriage, scrious infection, especially pelvic peritonitis, and severe anaemia from hacmorrhage are but some of the scars which a woman may carry for many ycars, as a result of such treatment. The scar of the psychological trauma may well be carried for life.

An attempt must obviously be made to put this deplorable situation to rights. Will the proposed reform of the abortion law do this?

Lord Silken's Bill, introduced in the last Parliament, emerged battered from its passage through the Lords, barcly recognisable through deletions and amendments only to dic a sudden death on the dissolution of Parliament. A modified version will again grind througl the legislative cogs of Westminster later this summer. If successful this will certainly clarify the legal position on abortion. It specifics who may perform such an abortion and in what circumstances. Thus an NHS gynaccologist, registrar or above - in agreement with the patient's G.P. - may terminate for the following reasons: if the mother's physical or mental health would be endangered by continuation of the pregnancy; if she were aged under 16 at conception, or mentally defective; or finally if the child would be likely to suffer from a defect which would prevent reasonable enjoyment of life. (Grounds of rape
and that the woman would make an inadequate mother were later deleted.)

This attempt at clarification is welcomed by many practitioners for whom the burden of decision is eased. Some, howerer, consider that the present law (which rests largely on the Bourne case judgement of 1938) allows considerably more freedom. Yet others fes that the change is not liberal enough and should take fuller account of social and cconomic factors as ground for termination.

In its present form the bill would do littl: towards climinating the criminal abortionists. Many of their patients are not those provided for by the bill but physically and mentally healtliy women, notably the single girl and the widow, whose pregnancy is looked upon with distaste by Socicty. For them the law will be effectively unchanged. For them the criminal abortionist will provide the only accepiable solution.

This situation could be improsed to some extent by liberalising the law - though not to the extent of "abortion on demand" as practiced in Japan and Czechoslovakia. The Swedish system of a pancl considering each casc on its merits, including socio-cconomic factors, has much to commend it.

An even more effective step would be widespread colucation in the most efficient methods of contraception. Yet cuen with better contraception mwanted pregnancies will occur. If we are cuer to be rid of criminal abortions and their dire sequalac, Socicty must view the unwelcome pregnancy through more sympathetic cycs.

## OVER-PRESCRIP'ГION

Overprescription of drugs has often been in the news in the light of the Annual Drug Bill, but recently another aspect has become prominent. It concerns the prescription of large quantitics of sedatives, anti-depressants and tranquilizers to a population which is increasingly employing them for self-poisoning rather than as remedial agents.

Unfortmately the people who are most likely to use these clrugs for sclf-poisoning are those to whom large quantities are given the depressed and the unstable. Surely it is time that other methods of making these drugs available were used. Kessel has already made this appeal in a recent article in the B.M.J. and Res Medica.

Could not greater control over prescriptions
be exerted? A weekly 'recurring' prescription requiring a weckly 'cancclling' signature from the pharmacist would limit the number of tablets issued at one time and incur no extra work for the Gl. The wider use of emetic charged burbiturates might be a worthy investmeant of the extra cost and more widespread dealing with the dangers of storing old tablets cie. would undoubtedly help. Some measures
might be more time consuming for the GP. in that he, as the 'family doctor' has the opportunity to wamn of the potential dangers of these drugs; put tablets into the custody of another member of the patient's family when necessary, and deal more thoroughly with psychiatric problems. But it remains the responsibility of the medical profession to consider priorities when discussing this problem.

## THE SOCIETY

Offec-bearers for the 23 oth session will be is. follows:
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'I'he Socicty's first year in llill Square has been cxtremely successful in both Public and Private busincss.

## PUBLIC BUSINESS

We have again been fortunate in laving many distinguished gucst speakers to address us. These included Professor F. J. Gillingham, Dr. W. I. Card, Dr. J. D. Roberston and Dr. Cicely Williams. Guest of honour at the President's dinner was Sir Dugald Baird whose address on some of the more amusing aspects of his carecr provided one of the highlights of the year. To these, as to all our other guests, we extend once more our thanks and appreciation.

## PRIVATE BUSINESS

The Private Business mectings have been described as the most important activity of the Socicty: this may or may not be truc, but certainly they provicle an opportunity for every member to be active in discussion. The value of this is twofold; it encourages members to learn to express their ideas in a more confident manner in public, and it is a stimulus to a wider interest in the art and science of medicinc.

The mectings of the past session were organised with this in mind. The majority of the meetings were introduced by a brief talk by a member and this was taken as the topic for discussion: topics ranged from "Prematurity" to "Exercise" and included many instructive clinical presentations. In each term two speakers were invited. In the first term Mr. J. Chalmers spoke on "Bone Growth" and Dr. R. A. Cumming on "The Blood Transfusion Scrvice"; and in the sccond term Dr. M. Gaze spoke on "Micro-clectrode Recording From The Hmman Brain" and Professor D. Whitteridge gave "Some Recollections Of Sir Charles Sherrington".

Essentially, however, Private Business mectings are what members make them: the more members that attend, the more members contribute, the more valuable are the meetings.

## REVISION OF LAWS

The Socicty's Laws have again undergone extensive revision by a committee set up for the purpose. The changes are concerned mainly with technicalities related to the clection of office-bearers. One welcome innoration, however, means that members may now entertain guests in the coffee lounge at any time.

# SEX <br> CHROMOSOME ABNORMALITIES IN THE MALE 

PATRICIA A. JACOBS<br>Medical Research Council, Clinical Effects of Radiation<br>Research Unit, Western General Hospital.

This article does not set out to give a compreliensive review of sex chromosome abnormalitics in Man, nor even in phenotypic males. Its purpose is more to outline a few gencral principles and slow how they apply to one group of individuals with one class of abnormality, namely males with abnormalities of number of either the X chromosome, or the $Y$ chromosome or of both.

Man has 46 chromosomes consisting of 22 pairs of autosomes, which are common to both sexes, and two sex chromosomes an X and a Y. Normal females have two X chromosomes which are morphologically indistinguishable from one another, while males have one $X$ chromosome and one $Y$ chromosome, which are morphologically dissimilar. (Fig. 1, Fig. 2). The main two critcria used in recognition of chromosomes are their length and the position of the centromere or primary constriction. On the basis of these criteria only 4 pairs of auto-
somes can be recognised with certainty, whilst the others can only be recognised as belonging to one of a number of groups. Some of these groups contain only two pairs of chromosomes, whilst the largest group, consisting of medium sized submetacentric chromosomes, contains as many as seven pairs of autosomes. Unfortunately the $X$ chromosome falls into this category of medium sized submetacentric chromosomes and, therefore, cannot be distinguished morphologically from autosome pairs 6-12. Nomal males, thercfore, have 15 chromosomes in this group - 14 autosomes and a single $X$ chromosome, while nomal females have $16-14$ autosomes and two $X$ chromosomes.

The Y chromosome, howerer, is one of the smallest chromosomes in the human complement with its centromere very near one end (acrocentric). It can usually be distinguished from the autosomes that it most closely resembles, those of pairs 21 and 22 , by virtue of
the fact that it never has small satellites on its short arms, and also because the constituent chromatids of the Y chromosome tend to be close to one another and are often rather fuzze: in appearance. Little difficulty, therefore, arises when dealing with abnormalities of mumber of the $Y$ chromosomes as these can be casily recognised. 'The X chromosome cannot. however, be distinguished morphologically from the autosome pairs 6-12. It is therefore necessary, when deciding whether or not one is clealing with an abnormal number of X chromosomes to consider, as well as the cytogenetic findings, the evidence from threc other sources, namely the muclear sex, the clinical findings and the results of autoradiography undertaken to determine the time at which the chromosomes syuthesise DNA.

Nuclear Scx
In a proportion of nuclei of non-dividing cells of normal females there is a small body about $1 \mu$ in diameter attached to the melear membrane. Such a body is nerer fomel in similar nuclei from normal males (Fig. 3). This body was first described by Barr and Bertram in $1949^{\text {a }}$ and is, therefore, referred to as the Barr body or sex chromatin body. Individuals having this body are said to be Barr positive or chromatin positive, whilst those in whom it is absent are said to be Barr negative or chromatin negative. There has been much sueculation since the first observation of sex chromatin as to its nature, but it is now known that it represents an X chromosome which is in a different state of condensation from the other chromosomes of the cell. While the latter are uncoiled and genctically active, the sex chromatin body consists of the whole, or the greater part, of an X chromosome which is condensed and presumably relatively inactive. It appears that for normal function the cell needs only onc X chromosome, and any further $X$ chromosome present is, to a greater on lesser extent, inactive, and represented by a sex chromatin body. Thas the maximum number of sex chromatin bodies is one less than the number of X chromosomes in the cell. Sex chromatin can be seen in the cells of a great many tissues in the body, but it is often studied in cells from the buccal mucosa, because these are very casily obtained. Such studies enable large populations to be sereened for their X chromosome status be means of a simple and quick techmique. The sex chro-
matin status of the individual gives valuable information as to the number and, in favourable circumstances, the size of the X chromosome, as abnormalitics of size of the X chromosome are reflected in abnomalities of size of the sex chromatin body. It must be emphasised, however, that it gives no information about the Y chromosome. 'I'hus indiv iduals who are lacking a sex chromosome, XO females, and also normal XY males are both chromatin negative, while both XXX females and males with Klinefelter's syndrome and an XXXY sex chromosome complement have two sex chromatin bodies in a proportion of their cells.


Fig. 1. Cell and Karyotype from a culture of the peripheral blood lymphocytes of a normal male.


Fig. 2. Cell and Karyotype from a culture of the peripheral blood lymphocytes of a normal female.

## Climical Features

Much work remains to be done on the clinical expression of abnomalities of the chromosomes in man. Yet it is clear that in gencral, abonomalities of the autosomes give rise to very much more severe and widespread anomalies than alonomalities of the $X$ and $Y$. fiuthermore where there are clinical manifestations of abnomalities of the sex chromosomes these tend to affect mainly the primary and secondary sexual derelopment, though they may also affect the mental development and belaviour of the affected individual. Thas the presence of an additional small antosome in
group 21-2 results in the profomed disturbmee of virtually every system in the body chatacteristic of mongolism. On the other hand the presence in an inclisidual of an additional Y' chromosome. which is of a comparable size. may be associated with no phonotypic abonormalities at all.

Fig. 3. Cell from the buccal mucosa of a normal female showing a single sex chromatin body (Barr body).

## Autoradiography

By trating cultures of homan cells with tritium labellect thymidine a specific radioactive precursor of DN: (luring the time the chromosomes are duplicating their material prion to division, it can be shown that there is a modium sizcel submetacentric chromosome in females. This suntlesises DND somewhet out of phase with all the other chromosomes in the cell, in that it both starts and finishes later. No such chromosome is present in cedls from normal males. 'This chromosome is presumed to be the "inactise" X which also forms the sex chromatin bodys and it has been shown that the mumber of late-syuthesising $\lambda$ chromosomes is one less than thie mmber of X chromosomes present in the ecell. 'I'o deter mine whether such a chromosome or chromesomes are present in the cell tritimo-labolled themidine is usually added to the cultures some three to four hous before haresting. when the majority of the chromosomes hate completed or are in the final stages of swo thesis. Single lavers of the cells are then pre pared on a microscope slicte and conered by a later of photographic emulsion which blackins when suitably exposed to a radio-active source. By this means ans chromosome, or chromosome region. which is actively syuthesising DNA later in the synthetic period can be recogniscel (l'ig. f).


Fig. 4. A normal female cell subsequent to treatment with trilium labelled thymidine which shows one very heavily labelled medium sized submeti:centric chromosome.

By way of illustration of how evidence from sex chromatin, clinical findings and autoradiography are correlated with the eytogenetic observations two cases can be considered. From cultures of both skin fibroblasts and peripheral blood leukocytes, both of these were shown to have fo chromosomes, the additional chromosome lecing a medimen sized submetacentric (hromosome indistinguishable from the chromosomes of group 6-12 . X. The first patient was a 28 vear old man who presented at a subfertility clinic. his wife having failed to conceive after 5 years of marriage. He was of werage intelligence and the only clinical featwe of note was the presence of small testes and azoospermia. The cells of his buccal mucosa were chromatin positive and autoradiography showed that he had a medimm-sized subuctacentric chromosomes which sunthesised DND late in the synthetic period. It was, therefore, concluded that the additional dromosome was and $X$ and the patient an example of Klinefelter's syndrome with an XXY
sea chromosome constitution. The second patient was an 8 year old mentally retarded boy who was noted at birth to have a cleft soft palate and who had subscquently been hospitalised on a number of occasions for severe respiratory infections. Ile also had a number of congenital anomalies, including hepomandibulosis, a sinus on bridge of his nose, abnomal dentition, curious facies and a systolic mumur. Repeated observations on cells from a number of tissucs showed him to be chromatin negative and autoradiography did not reveal any chromosome which swhthesised DNA out of phase with the others. It was, therefore, concluded that the additional chromosome was not an X chromosome, but was an antosome belonging to group ( $0-12$ and that the patient was trisomic for onc of these autosomes.

## Abnommalities of the Sex Chromosomes

The more commonly encounted numerical dbinomalities of the sex chromosome in males are listed in 'lable 1 , together with the sex chromatio and antoradiographical obscriations in these individuals. While these are the only abnomalitics which will be considered in the

TABLE 1

| Chromosome <br> Constitution | Sex <br> Chromátin | No. of Late <br> Synthesising <br> Medium Sized <br> Chromosomes |
| :---: | ---: | :---: |
| XXY | +ve | 1 |
| XXXY | ++ve | 2 |
| XXXXY | +++e | 3 |
| XXYY | +ve | 1 |
| XYY | $-v e$ | 0 |

paesent article it must be remembered that abnormalities of structure of both the $X$ and $Y$ chromosomes are encountered which may replace a normal sex chromosome or be additional to them. Furthermore it is very common to find, especially in individuals with abmonalities of the sex chromosomes, that not all the cells of the body have a miform constitution, but that thate are two or more cell lines present which can be distinguished cytogenctically, usually because they differ in their number of chromosomes. ()ne of the coll lines mas le nomal or all may be abnormal, and the resulting clinical picture depends on the constitution of the constituent cell lines and their relative frequency and distrib-
ution in the body, cspecially in the gonads. Thus an individual who has two cell lincs, one with 46 chromosomes and an XXY sex chromosome complement may, if his gonad is largely comprised of XY cells, be clinically indistinguishable from a normal male. Conversely, if his gonad is largely comprised of XXY cells he may be clinically indistinguishable from a "pure" XXY individual.

## XXY Males

The presence of a single cxtra $X$ chromosome is by far the commoncst abnommality of the sex chromosone complement found in man. This is associated with the features of seminiferous tubule dysgenesis or Klinefelter's syndrome. These features are somewhat variable but hypogonadism is always present. Before puberty, however, regressive changes take place characterised by marked degeneration and hyalinization of the seminiferous tubulcs, unusual numbers of Leydig cells and the absence of spermatogenesis. These features may be accompanied by abnormalitics of development of the secondary sex characters such as sparse growth of facial hair, female head or body hair distribution and development of breast tissuc. Furthermore XXY males tend to be rather tall and cmoichoid in proportion, their leg length being long in relation to their height. The I.Q. of males with an extra $X$ chromosome is on average lower than that of the normal population, such males being found significantly more often in institutions for the mentally defective than in the general population.

## XXXY

Males with two additional $X$ chromosomes are much less common than males with a single additional X chromosome. They also cxhibit the features of Klinefelter's syndrome but they are much more severely affected. Their testes are very small and the abnormalitics of development of the secondary sex characteristics arc usually very marked. Furthermore all XXXY individuals so far described have been mentally retarded - the majority of them being found amongst lower grade mental defectives.

## XXXXY

Males with three additional X chromosomes are, as expected, even more uncommon
and more severely affected than those with two additional X chromosomes. They also show a number of additional congenital abnormalities. Their testes are usually extremely small and often cannot be defined clinically. Some underdevelopment of the penis or scrotum is usually found, sometimes linked with hypospadias, and the degree of underdevelopment of the secondary sex characters is often very marked. There are usually marked skeletal abnormalities prosent, among them some degree of fusion or synostosis of the ulna and radius. All XXXXY individuals described have been low grade mental defectives.

## XXYY

Males with an additional X and an additional $Y$ chromosome are clinically similar to the XXY male. They have small testes and show a similar range of abnormalities of development of the scoondary sex characters. They are fairly uncommon, and all those so far described have been mentally defective. There is also some suggestion that they are taller than the XXY males and that they are musually prone to the development of acromegaly. XXYY males were also found to comprise one third of all chromatin positive males in an institution for criminal and hard to manage mental defectives - a proportion far higher than that found in ordinary mental defective institutions. It has therefore been suggested that the presence of an additional Y chromosome may be a predisposing factor to criminal behaviour ${ }^{\text {a }}$.

## XYY

Until recently relatively little was known about males with one additional $Y$ chromosome. Such individuals seem to be rare and the few cases described in the literature ranged from a normal fertile male cxamined because one of his children was a mongol, to a number of mentally retarded children with undescended testes. However, no clear picture of the XYY male had emerged, partly because they are indeed rather uncommon and partly because there is no casy way such as nuclear sexing, of recognising them in the population. Recently, however, a chromosome survey of the immates of a hospital for psychopathic criminals and for criminal and hard to manage mental defectives has been completed
and it was shown that 9 of the 314 men cxaminced lad an XYY sex chromosome constitutions. Clinical cxamination of the 9 XYY males showed them to be unremarkable, with apparently normal testes and genitalia." Howeser, they were significantly taller than the other males in the institution - in fact in this particular institution one in three of the males 6 ft . and over in height had an additional Y chromosome. While the frequency of XYY males in the general population is not known there is no doubt that their frequency in this particular group of patients is very much greator than could be cxpected by chance. This data suggests that the XYY male is a clinically unremarkable tall male, who is unusually predisposed to aggressive and criminal behaviour.

In conclusion it must be remembered that, while some of the more bizarre abnormalities of the sex chromosomes which have been described in this article are cxtremely rarc, chromatin positive males are common. Their incidence at birth is about 2 per thousand, which, if we assume there is no differential mortality, means that there are about 100,000 such individuals in the population of Britain at present. 'They form about $1 \%$ of all male mental defectives and have been shown to comprise over $10 \%$ of all azospermic and oligospermic
males attending a subfertility clinic ${ }^{\text {s }}$. Furthermore there is growing evidence that the prescnce of an additional $Y$ chromosome may contribute to psycopathic and criminal behaviour. It is, therefore, crident that males with abnomalities of the sex chromosome complement contribute very significantly to human pathology.

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# HEADACHE by Duncan L. Davidson, B.Sc. 

## From a dissertation read before the Society on February 4th, 1966.

Ilcadache is a symptom which may be a feature of a wide range of conditions, arising not only with pathology in the head but also in cardio-vascular, renal, metabolic, orthopacdic and psychiatric conditions. It is extremely common, and usually transitory. Yet it may be a symptom of great significance in clinical practice. The underlying mechanisms are largely unknown. The explanations of the cause of headaches have a long history, and it is on the aetiology of headache that this discussion will centre.

Ileadache can be a dramatic symptom and presumably has been ever since primitive man drank a fermented juice to excess and had his first hangover, or woke up with his first postconcussional headache to find that his wife had been kidnapped.
'Ihe first recorded description of headache comes from the 'Book of Prognoses' which is a series of tablets inscribed by the PhysicianPricsts of Mesopotamia probably before 2,500 B.C. These writings contain a mixture of clinical observations, which are often very astute, with statements on crude drug therapy, magic and religion.' One section on headache has been translated thus:
' when his brow pains a man and he romits and is sick, his eyes being inflamed, it is the hand of a ghost; then reduce to ashes human
bones and bray them; and anoint him with them in cedar oil and he will recover.'

It was scrious cnough having the "hand of a ghost' causing these afflictions but worse still
' not only the hand of a ghost but the hatred of a godeless against his life causes a man's right temple to hurt, his right cye to swell and tears to flow.'

While it is clear from the remains of neolithic skulls that trephing was performed at that time, and perhaps more surprising, that patients recovered, the precise indications for trephing are unknown. Perhaps an indirect indication may be considered from the finding that primitive groups in the South Sea Islands were trephing skulls during the last century. Possession by evil spirits was considered the cause of headache, epilcpsy and mental disorders; these spirits were released by trephining.
The contribution of Greck medicinc to our muderstanding of headache is the recognition that a particular group of clinical features constitute a distinct entity, the migraine syudrome. No advance in knowledge of the actiology of headache occurred with the Greeks, nor has it occurred until the present era.

Now, in our age of 'scientific enlightenment' we seek a more precise and objective description of the pathophysiological mechanisms than
demon possession, the hands of ghosts, and the hatred of goddesses.

In an article of this length the central mechanisns of pain perception cannot be discussed. Our concern here is with the disturbances which give rise to abnormal patterns of sensory input to the brain, and not with discussion of the fibres involved, the abnormal patterns themselves, and the central interprotation of these changes.

## Sources of Headaches

Knowledge of which structures in the head may give rise to pain comes from two main sources. The first of these is the correlation of clinical obscrvations, c.g. an occipital headache, with pathological findings, c.g. an infratentorial tumour. The second source is from neuro-surgical operations under local antacsthesia in which records are made of the site of referred pain which is clicited by such crude methods as crushing, stretching, distending, burning and electrically stimulating various structures of the head.t There is, however, a great difference between cliciting pain in an operation and demonstrating the mechanisms which operate in the clinically occurring headache. The following results, therefore, arc but crucle pointers to the sites at which pain may arise. The scalp, as expected, is pain-sensitive while the cranial bones are insensitive. The dura appears to be insensitive to pain-producing stimuli except in the vicinity of arteries, venous sinuses and their tributary veins, and the floor of the anterior and posterior fossac. Pain is particularly easily elicited from the large arteries at the base of the brain. Cranial nerves V, VII, IX and X, all nerves with a sensory component, and also $\mathrm{CI}, 2$ and 3 are pain sensitive. No pain was clicited from the pia and arachnoid mater, the parenchyma of the brain or the linings of the ventricles. As the supra-tentorial dura is innervated by branches from the Trigcminal Nerve, pain arising in this region is usually referred in the distribution of the nerse producing peri-orbital, frontal and temporal headaches. Infra-tentoral disease, on the other hand, is largely referred via Cranial nerves IX and $X$ to the auricular region and via the upper cervical nerves to the occipital region and the upper part of the neck.

The causes of headache have, incvitably, been classified. One simple and useful approach is to consider that headaches may arisc in any, or a combination of five general wavs ;
(1) traction upon intra-cranial structures
(2) intra-cranial inflammation
(3) Vascular changes
(4) sustained contraction of scalp and neek muscles
(5) spread of pain from discases of the eyes, cars, nose and throat.

## Traction Upon Intra-Cramial Structures

Headache may occur with expanding intracranial lesions whether these be neoplasms, subdural or intra-cerebral haematomas, or abscesses. It is almost always a presenting symptom in infra-tentorial lesions presumably because expansion occurs within a confined space in a region which contains a number of pain-sensitive structures. With expansion of the lesion distortion of the normal anatomy occurs, and pain arises from traction on and displacement of nerves and vessels. In supratentorial lesions headache occurs as a presenting feature in only about a third of all cases, presumbaly because greater expansion may occur before pressure and traction effects become prominent.

## Intra-Cramial Inflammation

The intra-cranial inflammation that occurs in meningitis or sub-arachnoid haemorrhage is associated with severe headaches. The pain probably arises in part from vascular changes presently to be discussed, partly from traction and pressure effects, and perhaps from direct stimulation of nerves by the ill-defined entity 'toxins' and the breakdown products of affected cells.

## Discases of Eycs, Ears, Nose, Throat and Tecth

Discases of the cyes and orbit, E.N.T. conditions like sinusitis and acute otitis media, as well as dental abscesses may produce pain. In gencral the pain is at first localised at the site of the lesions, but with progress of the disease process it may radiate in the distribution of the nerve involved.

## Headaches of Vascular Origin

The vascular changes which occur in a number of conditions, for example, hypertension, migraine, uraemia, and febrile illnesses appear to be related to the headaches. The subject is perhaps best approached through an experimental model, the headache induced by the intravenous injection of histamine. Within a fow seconds of injecting histamine there is a flushing of the skin, hypotension and a rise in C.S.F. pressure. In about 30 seconds the blood and the C.S.F. pressures return to nor-
mal, and it is at this time that the headache begins. It is a bilateral, throbbing headache whech usually lasts between 10 and 20 minutes. 'Ihere is only indirect evidence that the headache is related to intracranial vasodilatation and the evidence is as follows. 'I'he oscillations in C.S.F. pressure that are in phase with the arterial pulse are increased during the headache. 'ithe hoadaches are reduced by mancourres which reduce the intracranial arterial pressure. 'The headaches are intensified or diminished by lowering or clevating the C.S.F. pressure respectively.

Wolff has applied these methods to patients with headaches and finds that a number of headaches are altered by these manocurres in a similar mamer to the histamine induced headaches. This group includes the headaches of uraemia, all the febrile illuesses, post-scizure and post-concussional (in part). As vasodilatation is common to all these states it is commonly stated that dilatation of the ressels, perhaps with stretch of the fine nerve endings in the wall, is the cause of the headache. But it is an inadequate explanation, for it does not recognise that there may be a process which has in common rasodilatation and pain stimulation. Recent work on the headaches of the migraine syndrome suggests that, in migraine, arterial dilatation is only part of the story. Ancl so it may be revcaled with further investigation that the headaches due to intracranial vascular changes have mechanisms similar to that in migrainc.

The headache in the migraine syndrome is classically a unilateral throbbing headache that may be peri-orbital, frontal, temporal, or occipital. It may last from under half an hour to several hours. But the heaclache is only part of a syndrome which may be very variable in presentation. In about $15 \%$ of patients there may be prodromal symptoms occurring between 20 and 40 minutes before the onset of the headache. These may include a varicty of visual changes, such as scintillations, scotomas, or even hemianopia. There may be paracsthesia, ataxia, vertigo, or changes in consciousness or mood.

But what is known of the underlying mechanisms in this condition? When the vessels of the bulbar conjunctiva are directly examined and photographed, arteriolar constriction is found in the prodromal phase. The finding that E.E.G. changes are consistent with focal cerebral ischaemia, and that the prodromata can be reduced or abolished with breathing $10 \% \mathrm{CO}_{2}$ mixtures suggests that arteriolar
constriction with areas of cercbral ischacmia may underlic the prodromal phase.
the origin of the headache of migraine in the extra-cranial arterics is suggested by a number of observations. Pain can be reduced or abolished by dircet pressure or procainisation of the extra-cranial arteries. Unlike the histamine induced headaches it is maffected by manocustes which alter intra-cranial pressure. During a headache the superficial vessels beconce tender, painful and surrounded by ocdema fluid. Simple measurements with a tambour show increased amplitude of pulsations during the headache. Three changes must be explained during the headache phase; (1) vasodilatation, (2) oedema formation, (3) pain production. What may be implicated as the perpetrator of these changes.

The relcase of endogenous histamine from a bound form has been considered an unlikely mechanism as the headache differs in its characteristics from a histamine induced headache. Also, the migraine headache is unaffected by anti-histamines in contrast to the reduction in pain that is produced in the rarer, rather bizarre condition 'histaminic cophalgia' or 'cluster headaches'. Recently, however, Schayer' using radioactive tracer techniques, has shown the existence of an 'induced' form of histamine, that is maffected by anti-histamines, and has a longer time course of action than 'bound' histamine. Its precise role is as yet speculative. But histamine cannot yet be dismissed as having no role in the headaches of migraine.

May 5 H.T. or a kinin be a cause of the vascular changes? To investigate this, small quantities of tissue fluid have been aspirated from the vicinity of the temporal arterics during and immediately after the migraine headaches and also in headache-frec intervals. These were then compared with the aspirates from nomal subjects by a number of phamacological assay methods.

Chapman' claims to have found in the tissuc fluid a polypeptide and also an enzyme that, on incubation with plasma, is capable of producing increased quantities of the polypeptide. This polypeptide is similar but not identical to bradykinin, and clearly differs from 5 II.T. The levels during the headache period were on average about 8 times that of normal subjects, and it was stated that the level correlated very well with clinically estimated sererity of the headache. Here, of course, is an illustration of one particular difficulty in work on pain - that of estimating the severity of pain and comparing the severity of pain in one person with that
in another. This polypeptide has been labelled 'neurokinin'. While it does fulfil our three criteria, those of rasodilatation, increased permeability with ocdema formation, and pain production, this work is rather tentative. It may be that the improved methods of separation and characterisation of the kinins involved will throw more light on the subject. It may become crident that a multi-factorial mochanism is responsible for the headache. It must also be pointed out that no progress has been made on the questions of what initiates the whole process, why the headache should be unilateral, and why the changes should occur only in the external carotid vessels? The autonomic changes which may occur in association with the headache are varied; there may be lacrimation, bradycardia, sweating, nasal congestion, constipation (or even diarrhoea). It may ceentually be shown that the migraine syndrome takes origin in a disturbance of autonomic function.

The headaches of hypertension, on the same evidence as in migrainc, appear to originate in changes in the extra-cranial arterics. The headaches do not, however, relate directly to the level of the blood pressure except during acute attacks. The fact that the arterial pressure is clevated throughout the body, but only the extra-cranial vessels become painful suggests that there may be some intrinsic difference in the extra-cranial arteries that renders them liable to develop pain. Little work has been done on the lieadache in hypertension so that it is only an interesting speculation that the mechanisms may be similar to those of migraine.

## Muscle-tension Headachc

The other common chronic headache, a cause of distress to thousands of patients, is one which often arises in relation to stress or anxicty and is called a 'tension' or 'muscletension' headache. Its characteristics are that it is non-pulsatile, fairly constant in intensity, being a dull ache rather than an acute pain. Patients describe it rariously as a 'tight band' or cap or as an oppressive pain on the top of the head. Often it is bilateral in the occipital region, extending into the nock, but it may vary and be cither umilateral or bilateral, and may occur in the frontal or parictal arcas. It
may occur primarily in a neurotic type of reaction or in anxicty producing situations, but it may arise sccondary to pain elsewhere, e.g. migrainc, or secondary to degenerative changes in the cervical spinc.
'The term 'muscle-tension' hoadache has evolved because one feature of the hadache is a sustamed contraction of the occipitofrontalis, temporalis or the muscles of the neck. E.M.G. recordings from these museles showed increased activity during the headache periods which was approximately related to the 'clinical anxicty level' (again a very subjective and unreliable estimation). There is, however, another factor besides increascel muscular actisity, and this is the observation that arteriolar constriction occurs in the bulbar conjunctive during the headaches. As the headaches are intensified by vasoconstrictors and relieved by vasodilator drugs it is reasonable to infer that constriction may be occurring in the ressels of the active muscles and that this contributes to the headache. Presumably, thercfore, headache arises from a combination of ischaemia due to vasoconstriction and increased metabolism of sustained muscular contraction.

## Conclusion

Headaches are often trivial and transitory and as such tend to be ignored climically. But they arc important in two ways. Firstly, the chronic, recurring headaches cause considerable miscry in the community. Sccondly, headaches may be of great clinical significance, for example in hypertension and in intra-cranial tumours. The explanation of the causes of these headaches have varied through the ages, and cven now ideas change as information increases. Some current ideas have been outlined in this article. As knowledge increases in the future so should the treatment of the underlying causes improve.

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## BOOK

## REVIEWS

CALLING THE LABORATORY. Editor: W. A. R. Thompson, M.D. 2nd edition. E. \& S. Livingstone, Ltd. 1966. 17/6d. pp. 126.
This book, although designed for use by general practitioners, will fill a long-felt need of both students and hospital staff.

It gives a brief outline of the various laboratory tests commonly in use today, with an explanation of the indications for such tests, and the principles and methods underlying them. Perhaps most useful of all, and certainly most practical, it tells doctors the correct way in which to present the specimen under test to the laboratory - surely a sore bone of contention between doctor and laboratory for many years.

As a practical review of laboratory tests it is admirable, and fully recommended. Its title and cover illustration, however, could well be improved.
D.B.

## GLAUCOMA: EPIDEMIOLOGY, DIAGNOSIS

 AND SOME ASPECTS OF TREATMENT. Proceedings of a symposium held at the Royal College of Surgeons of England 1965.The practical difficulties in mass screening of the population to detect unsuspected glaucoma in its early stages are enormous. This is especially so if one realises that "suspects", when discovered, constitute about $10 \%$ of the population and that further investigation of these will lead to discovery of the disease in about $1 \%$. Unfortunately the problem does not end there.

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Not a recommended buy for the student unless he, for better or worse, wishes to complicate his thoughts on glaucome.
B.C.

TEXTBOOK OF MEDICAL TREATMENT. Edited by Sir Derrick Dunlop and Stanley Alstead. 10th edition. E. \& S. Livingstone, Ltd. 1966. 70s. pp. 1003.
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I.C.M.

A GUIDE TO CARDIOLOGY. By J. C. Leonard and E. G. Galea. 2nd edition. E. \& S. Livingstone, Ltd. 1966. pp. 306.

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