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GIANT-CELL GRANULOMA OF THE RESPIRATORY TRACT (WEGENER'S GRANULOMATOSIS)

BY

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This paper is based on a retrospective study of data from 10 patients who had in common an illness characterized by symptoms of progressive ulceration in the respiratory tract together with signs of widespread inflammatory disease. Histological examination of material from each case shows disseminated granulomata, most common in the respiratory tract and kidneys, and widespread vascular lesions similar to polyarteritis nodosa. The name Wegener's granulomatosis has been applied to this syndrome (Ringertz, 1947; Johnsson, 1948; Fahey et al., 1954). The purpose of this paper is to describe in brief the clinical and pathological features, to give a concept of the pathogenesis, and to suggest a method of treatment. The cases, which were selected according to the pathological criteria of Godman and Churg (1954), have been described in detail elsewhere (Leggat and Walton, 1956; Walton, 1957).

Clinical Features

The main data from each case, and from 46 others selected from the literature, are summarized in Table I. An analysis of symptoms and signs is given in Table II. Typically, the disease occurs in previously healthy young or middle-aged adults of either sex. The onset is insidious, with non-specific symptoms of infection in some part of the respiratory tract. Two patterns can be distinguished. In about two-thirds of the cases persistent purulent rhinorrhoea is accompanied by nasal obstruction and crusting, antral pain, and epistaxis. Otorrhoea, deafness, or ulceration of the gums was the initial symptom in a few of these: each later developed rhinorrhoea. In the second, smaller group, attention is drawn to the lungs because of chronic cough, haemoptysis, or pleurisy. Often the constitutional upset is out of proportion to the apparent intensity of the local lesion, and the patient seeks advice because of persisting malaise, fever, or weakness.

The course is usually rapid and full of incident, progressing to death in, on average, five months, occasionally in as little as four weeks. A few patients (Cases 9, 19, 43, 45, and 47, Table I) have had a more chronic illness with periods of remission, and survival for up to four years. Though temporary improvement sometimes follows antibiotic treatment, the local lesion always persists. Spread of the inflammatory process leads to extensive mucosal ulceration and cartilaginous or osseous destruction in the nose and palate on the one hand, and to widespread pulmonary consolidation on the other. Spread through the upper air passages is often followed by conjunctivitis, dimness of vision, increased lacrimation, and exophthalmos, deafness, earache, and otorrhoea. The development of a sore mouth, hoarseness, or dysphagia has resulted in the discovery of ulceration in the fauces, pharynx, or larynx. Only twice (Cases 1 and 53) has the mucosal ulceration spread to involve the skin of the face.

Sooner or later signs of widespread inflammatory disease appear in every case: fleeting arthralgia, numbness and tingling in the limbs, sensory loss, muscle weakness or paralysis, and a haemorrhagic vesicular rash most frequent on the skin of the face, wrists, and elbows and the oral mucosa are all common. A pericardial friction rub or electrocardiographic changes occasionally indicate involvement of the heart; parotitis, orchitis, and prostatitis also occur. In the late stages fever, usually of septic type, is almost constant, and albuminuria, haematuria, cylinduria, and/or pyuria have indicated renal involvement in nearly every case.



FIG. 1.-Case 10. Multiple rounded opacities are present in the left upper zone and both lower zones. Spicules of calcification are seen in the left upper zone.

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Cause of Death		Respiratory failure ? Uraemia Respiratory failure	Uraemia Uraemia Presinatory failure	I Iraamia	Oracinia (Alive)	? Úraemia	Uraemia	Uraemia		Cardiac failure	Uraomia Cardiac failuro	Respiratory failure	Sopsis, cacricata	Uraemia Respiratory failure	I [reemia		Respiratory failure	Uraemia	Cardiac failure	Uraemia	Respiratory failure Uraamia	::	Septicaemia	Respiratory failure	Uraemia 	Cardiac failure	Uracmia	Myocardial infarction	Myocardial infarction	Cardiac failure	Cardiac failure
USUSCO LINGLA		+ 1 + -		+ 4	++	+	i + +	+++	+	-+	++1	+-	+ •	+++	+++	+ +	+++	+	+ +	1+	++	++	++	++	++	+	++	1+	/ <u> </u>	+	++
LSSU2	10	1.1.1		11	+	11	11		I	1	1+1	1		111	111			t	+1	11	11	1+	+1	j I	11	1	11	11	ł	1	11
omobathy	- N	+11-	FII	1	1	11	11		+	-+	+11	+	I	111	111	-	F11	ł	1+	+1	11	1+	+ 1	11	+1	+	1+	+1	I	1	1+
ini eagust	CP	+-	+ 1 +	- 1 1	+	+	++	111	1	1	+	1	I	111	111	I	1+1	1	+1	1+	+1	++	• 1 +	+ 1	++	1	+ 1	1+	+	+	11
dzaA ni	۶R	+++•	+11	14	++-	+ 1 -	++-	+	I	I	+	.+	I	11+	111	1	11+	1	+1	11	11	++	+ 1	+ 1	1+	1	++	1+	+	+	+1
r Changes	Ea	1++	+ 1 -	+ 1	11	11	++		4	F I	111	I	1	11+	1+1	1	1+1	ł	11	+1	11	11	1+	+		I	+1	1+	1	1	1+
e Changes	EY	+ + + + •	+		1+	+	+ 1)		+	+	ł	1	111	+ +	F	1+1	Ì	++	1+	- 1 1	+1	+1	++	+ 1 +	1	++	·++	1	+	+1
t Symptoms	Lower	Cough Nii Stridor, dyspnoea	Dysphoea	Pain in chest	Nu Haemoptysis	Productive cough Nii		Productive cough Nil Chest pain	Nil	Productive cough	Purulent sputum Productive cough Nil	Cough, haemoptysis	Nil	Productive cough Nil Chest pain. cough	NII		Productive cougn Nil " "	Pain, dyspnoca	Pneumonia Nil	Cough Pain hemontuele	Cough, haemoptysis	Nil Couch chest pain	Nil	Dyspnoea	Chest pain Nil		: : :	:::	•	:	Cough, pain Cough
Respiratory Trac	Upper	Epistaxis, rhinorrhoea Nasal block, epistaxis Purulent rhinitis	Anosmia, oral ulceration Nasal block, rhinitis	Sore mouth •	Hoarseness, sore throat	Epistaxis Purulent rhinitis	Rhinitis, dysphagia Saddle nose, hoarsences	Foul rhinorrhoea Fetor, gum bleeding Headache	Titoauacue	Earache Epistaxis	Sanious rhinitis, stridor Sanious rhinitis Frittavis masal block	EN	Nasal obstruction	Nil Otorrho ca Fristaris, otorrho c a	Sanious rhinitis Earache	Gingrvitts	Nil Nasal obstruction Epistaxis, aphonia,	retor Nil	Crusting and hoarseness Bpistaxis	Epistaxis, hoarsenees	Distants, sconstitutes	Epistaxis Cont Iconstice	Rhinorrhoea	Purulent rhinitis	Saddle nose Purulent rhinitis Oral ulceration	Nasal deformity	Sinusitis Duenheorie	Saddle nose Purulent rhinitis	Loosening of testh	Sinusitis	Nasai block, hoarseness Otorrhoea
Initial Symptoms		Sore gums, limb pains Otorrhoea, nasal obstruction Nasal block, joint pains	Pleurisy Dyspnoea	Deafness, sinusitis Cough, dyspnoea	Malaise Cough, joint pains	Rhinitis with obstruction Sinusitis	Joint pains, septic illness Jaw pain, rhinorrhoea	Rhinorrhoea Sanious rhinitis, headache	KIMOTTOCA, OPEIAAIS	Otorrhoea Cough	Dyspnoea, dysphagia Saddle nose, epistaxis	Leg pains, sore throat	Gum ulcer, cough	Cough, fever Deafness, earache Sinnisis	Sinusitis Rhinorrhoea, headache	Rhinitis with obstruction	Cough with sputum Deatness and headache Nasal obstruction	Chest pain, dyspnoes	Sinusitis Rhinitis with obstruction		Hadmoptysis, cnest pain Dyspnoea, cough	Sinusitis, archectasis	Orcantus, weakinces Sinusitis	Sinusitis, otica Rhinorrhoea, sinusitis	Sinusitis, cough, pleurisy Sinusitis, fever Senious rhinorrhoes		Blooding from our	Coryza, hoarseness Earache	Gingivitis, neck ulcer	Epistaxis, weight loss	Chest pain, cough Fever, earache
Duration Months			0 m	م ٹ	21+	12	ي. •	<u>ب</u> ممر	•	~~~~~	90 11 30	۹ I	∞		040	\$	2 4 3	7	40	9	-=-	n 7 .		*	804	2 5	3	144	Ħ	•	-0
Onset		1943 1948 1948	266 726	1954 1955	1947	1955	1691	566 566 566	6661	1940	1938 1939		1945	1947	132	6461	8666 8666 8676	1948	1948 1951	1951	2020	561	266	1953	1951		2	1955	1953	1955	1950
Age and Sex		8833 MM	42 7 1	22 22 22	28 MM	55 F	38 22	33 F	1	32 F	868 744	2 P. 2 9	37 1 1	285 7 F	222 227	23 M	r⊼X 8 ≌	70 M	4€ ₽₩	51 M	NN NN	9%: 9%:	22; 88;	4 7 7 8 7 8	91 38 31 38	2 2	1 2 2 2 2 2		26 M	42 M	26 27
Description		Wegener's granulomatosis				Periorianitie scoloss	Rheumatism Rhinogenic granulomatosis		Periarteritis nodosa with lung changes	Periarteritis nodosa	Sarcoidosis Chronic graauloma	Unknown granuoma and per- arteritis nodosa	Periarteritis nodosa; an unusual	Vegener's granulomatosis	Lettal muune granuloma Malignant granuloma	Giant-cell granuloma	Complications of ainusitis Idiopathic nasal granuloma	Respirato-remal type of periarteritis	nodosa. Nasal rianuloma and periarteritis	nodosa ,,	Necrotizing granulomatosis of lung	Wegener's granulomatosis	::		Nasal granuloma, periarteritis	arteritis nodosa		Utant-cell gramuoma Wegener's gramulomatosis		Wegener's anglitis	Stevens-Johnson syndrome Polyarteritis nodosa
Author		Walton	:	:	:	V15000 (1031)	Rössle, 4 (1933) Wesener (1936)	: :	Postel and Laas (1941)	Banowitch et al.,	2 (1942) Staehelin (1942) Lindsay et al. (1944)	Wonderg (1946)	Ringertz (1947)	Johnsson (1948)	Wullams, 3 (1949) McCart (1950)	Howells and Eried-	Mallory (1950) Pugh (1950) Woodburn and	Ahlström et al.	(1953) Stratton et al.	(1953)	Ficaberg (1953a)	Fahey et al. (1954)	::	::	McCallum (1954)	cox (1954)		Milner (1935) Cogan, 2 (1955) Bienami and Ficari	(1956) Morgan and O'Neil	(1956) Chatillon et al., 3	(1956) Short, I (1957) Rose and Spencer, 87 (1957)
		-44	45	50	~ 0	2=	122	22	16	18	 <u></u>	2 2	ង	222	223	ล	823	33	23	8	333	84	44	44	3 41	•	\$ \$	828	3	3	22

TABLE I.—Clinical Features of 56 Cases of Wegener's Granulomatosis

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BRITISH MEDICAL JOURNAL Once signs of dissemination have occurred the general condition has always deteriorated and death has ensued from uraemia or secondary bronchopneumonia (Table I).

Clinical investigations of note are summarized in Table III. The radiological changes in the lungs are of particular interest. In just over half of the cases

TABLE	II.—Symptoms	and	Signs	in	56	Cases	of	W	egener	's
	• •	Guan	wloma	tre	i.					

								Cont.
			••	50	•••		89.3	
or sinu	ısitis	••	••		34	••		60·7
witho	ut rh	inorrho)ea		- 4	••		7.1
••	••	••	••		18	••		32.1
••	••	••	••		5			8.9
R.	••	••			7	••		12.5
••	••	••	••		8			15-3
••					3			5.4
deafn	CSS .	••	••		18			32.1
				23			41.1	
••	••	••	••		17			30.4
•••	••	••	••		-5			8.9
	••	••			3			5.4
					6			10.7
••	••	•••		07			40.0	
••	••	••	••	21	22	••	40.7	20.2
••	••	••	••		4	••		39.3
••	••	••	••		11	••		10.6
••	••	••	••		11	••		19.0
••				- 14		••	25.0	
••					10	••		17.9
••					4	••		7.1
••	••			26			46-4	
••		••		19		••	33-9	
••	••	••	••	16			28.6	
	(or similar without a similar simi	or sinusitis without rh: 	or sinusus	or sinusits	or sinusitis	for sinusitis	for sinusitis	or sinustics

I ABLE	III.—Clinical	Investigations	<i>in</i> 36	Cases	OJ.	wegener's
		Granulomat	osis			

	No. of	Abn	ormal
	Examined	No.	Percentage
Radiography: Skull: sinus opacities Lung shadows . Round and discrete Bronchopneumoaic infiltra- tion	21 40	18 38 23 10	85-7 95-0 57-5 25-0
Haematology: Microcytic anaemia Leucocytosis Eosiaophil leucocytosis	33 56 37	32 38 17	97·0 67·9 45·9
Urine: Albunin Erythrocytes Leucoytes Casts	42 42 42 42 42	37 34 21 28	88-0 81-0 50-0 66-7
Biochemistry: Uraemia Hyperglobulinaemia	27 10	23 8	83·2 80-0
High blood pressure	33	8	24.2

dense circular or oval opacities are present in one or more lobes (Fig. 1), often showing central cavitation, and varying in size up to that of a hen's egg. Less often bronchopneumonic infiltration is the only change of note.

No specific aetiological agent has been isolated in any case. From examination of sputum and nasal swabs in 24 cases, the only pathogens isolated were *Staphylo*coccus aureus from 14, a haemolytic streptococcus from 7, Streptococcus viridans from 4, and pneumococci from 3. Culture for fungi was negative in five out of eight cases; in the others Actinomyces bovis (twice), Candida albicans, and a "leptothrix" were isolated. A positive blood culture, a positive blood Wassermann seaction, and isolation of tubercle bacilli have each been reported once, but otherwise these investigations have always been negative. Similarly, all agglutination tests have been negative.

Pathology

The most striking changes occur in the respiratory tract. Extensive ulceration, often widespread, with necrosis of underlying cartilage and bone, is present in the nose, paranasal sinuses, palate, pharynx, and/or glottis in about threequarters of the cases (Table IV). Most also show multiple

 TABLE IV.—Sites of Respiratory Tract Ulceration in 56 Cases of Wegener's Granulomatosis

					No.	Per Cent.
Nose and :	nasal si	inuses			34	 60.7
Mouth and	d phary	/mx			21	 37.5
Tongue					7	 12.5
Larvnx					14	 25.0
Traches					17	 31-5*
Bronchi					23	 42.6*
		* Per c	ent. of	54 ca	ses.	

shallow ulcers in the trachea and bronchi. The lungs were abnormal in all but two cases (Nos. 20 and 50). Rounded nodular areas of consolidation, greyish white in colour, measuring up to 4 cm. in diameter, and sometimes showing central cavitation, are usually present in one or both organs. Less frequently pale miliary nodules scattered diffusely through all zones, or varying degrees of consolidation, are noted. Infarcts were described in Case 36 only.



F10. 2.—Case 3. Lung. (Haematoxylin and eosin. $\times 65$.) A bronchiole showing intense granulomatous inflammation with early necrosis. Two giant cells of foreign-body type are present.



Fis. 3.—Case 7. Lung. (Haematoxylin and eosin. ×230.) Granuloma with numerous giant cells and extensive central necrosis.

giant cells with a sprinkling of lymphocytes,

cells, and both neutrophil and eosinophil

morphonuclear leucocytes. Central necrosis of mature lesions is constant. The vascular les-

ions are focal, in-

dependent of the

granulomata, and

affect small arteries

and arterioles, oc-

casionally venules.

The acute stages

(Fig. 4) show "fi-

brinoid" necrosis

of all layers, des-truction of the

muscle and elas-

tica, mural thromb-

osis, and neutro-

plasma

poly-

Irrespective of their site, the lesions of the respiratory tract in my cases have a constant histological pattern (Fig. 2), that of granulomatous inflammation spreading deeply from the mucosa of the respiratory passages into the surrounding Intrapulmonary bronchial ulceration is especially tissue. common. Giant cells, of both Langhans and foreign-body type, are very numerous in the granulation tissue, and widespread necrosis often affects both cartilage and bone. Many of the pulmonary lesions are clearly peribronchiolar or prove so to be in serial section studies, and the rounded areas of consolidation visible macroscopically are seen to be conglomerate areas of peribronchial and peribronchiolar Vessels near the lesions are implicated in the necrosis. inflammation but show no features to suggest a specific vascular disease: thus the changes in these vessels are often confined to the segment next to the granulomatous focus. No micro-organisms can be demonstrated in appropriately stained sections of lesions in any of my cases.

In addition to the ulceration of the respiratory tract, widespread granulomata and foci of vascular necrosis are present in each case. Most often perivascular, the granulomata are discrete, of constant tuberculoid pattern (Fig. 3), and consist of endothelioid cells in radial arrangement and Langhans type



FIG. 4.—Case 4. Spleen. (Haematoxylin and eosin. × 90.) Pulp arteriole at a bifurcation. One limb shows acute "fibrinoid" necrosis and perivascular polymorph infiltration. Fig. 4 -

phil infiltration. Healing lesions, present in most cases, show organization and recanalization of both the necrotic vessel wall and the luminal thrombus.

These widespread lesions, the incidence of which is given in Table V, are most frequent in lung, spleen, and kidney, though the number varies from case to case and organ to organ. Sometimes-for example, Cases 8 and 19-the granulomata are very numerous and visible as pale nodules about 0.1 cm. in diameter in many organs; in others the vascular lesions are very numerous, producing infarcts in kidney, spleen, heart, and testis.

The kidneys deserve special mention, having been severely affected in all but Cases 18, 30, and 45. Enlargement and blurring of the architecture are the usual macroscopic changes. Focal "fibrinoid" necrosis of glomerular loops and periglomerular granulomatous inflammation, often with giant cells, are almost constant histological findings. In my cases about a third of the glomeruli are usually affected, the inflammation and necrosis frequently involving the afferent or efferent arterioles.

Discussion

Despite the wide range of symptoms occurring in these cases, a constant clinical pattern is clear-that of a disease process beginning with symptoms and signs of an inflammatory lesion in the respiratory tract, continuing with evidence of widespread inflammation, and terminating in renal or respiratory failure. In most, especially Cases 4, 9, 10, 47, 48, and 49, the priority of the lesions of the respiratory tract is quite clear, and a definite time interval, of up to three years (Case 47), has often elapsed before the onset of widespread disease. In the early reports (Klinger, 1931; Rössle, 1933; Wegener, 1936, 1939) nasal lesions were prominent. Later authors (Postel and Laas, 1941; Weinberg, 1946; Ringertz, 1947; Johnsson, 1948; Godman and Churg,

TABLE V.-Incidence of Disseminated Lesions in 54 Cases of Wegener's Granulomatosis

				No.		Per Cent.
Discrete granuloma	ta		• •	54		100
Upper air passag	65			28	•••	51-9
Trachea				10		18-5
Lungs				44		81-5
Spleen				30		55-6
Kidney				36		66.7
Liver				ġ		16.7
Lymph nodes			•••	ú		20.4
Heart	••	••	••	6 -	••	11.i
Prostate	••	••	••	Å	••	7.4
Brain and mening		••	••	7	••	7.4
Other organs		••	••	Å	••	7.4
Other Organia	••	••	••	-	••	1.4
Focal necrotizing as	rtari	olitis		54	·	100
Unner air passag	AR		••	14	••	25.9
Lungs		••	••	47	••	87.0
Spleen	••	••	••	42	••	77.9
Kidney	••		••	42	••	77.9
Liver	••	••	••	76	••	19.5
Lover	••	••	••	16	••	10.7
Intestine	••	••	••	13	••	21.0
	••	••	•• •	13	••	44.1
	••	••	••	~	••	10.7
Adipose dissue	••	••	••	0	••	, 14.0
voluntary muscle	•	••	••	8	••	14.9
Pancreas	••	• ••	••	. 3	••	9.3
Aurenal	••	••	••	6	••	11-1
lestis	••	••	••	5	•••	9.3
Gail-bladder	••	••	••	4	••	7.4
Peripheral nerve	••	••	••	- 4	••	7.4

1954), in describing cases in which tracheo-bronchial and pulmonary changes were more striking, noted their simlarity to Wegener's cases. As each type shows similar local and widespread histological features, it is clear that they differ only in the level in the respiratory tract of the initial lesion. Furthermore, tracheo-bronchial or pulmonary lesions were present in all except Cases 20 and 50, and lesions in upper air passages may have been overlooked in some or all of the 12 cases in which such were not recorded.

To date, including those of the present series, at least 56 cases which satisfy the diagnostic criteria of Godman and Churg (1954) have been reported, often as polyarteritis nodosa, malignant granuloma, or lethal midline granuloma. There can be little doubt, though, that the condition is more common than at first appears, as many cases have been excluded from the series only because of the absence of full data-namely, those of Siegmund (1936), Schürmann (1936), Moore et al. (1951), Rojas (1952), Geist and Mullen (Case 1, 1953), Breckenridge et al. (Case 3, 1954), Alexander (Case 3, 1954), Cutler (Case 2, 1955), Friedmann (Cases 4, 5, and 7, 1955), Paterson (Cases 1, 2, and 6, 1956), Rogers and Roberto (Cases 1 and 2, 1956), French and Civin (1956), Kelly (1956), McKibben and Bayliss (1956), Chatillon et al. (Cases 1 and 2, 1956), and Singh et al. (Cases 1 and 4, 1958), and others mentioned by Godman and Churg (1954). A further estimate of the frequency of the condition can be gained from the analysis of 104 cases of polyarteritis nodosa by Rose and Spencer (1957). Of these, one, Case 87, is a typical example of Wegener's granulomatosis, while four others, Cases 81, 82, 84, and 89, had granulomata of the upper respiratory tract and are almost certainly further examples.

Of the 56 cases, 33 have been male and 23 female. All age groups save the very young have been affected, the youngest patient being aged 12 years, the oldest 75, with the peak incidence in the fourth and fifth decades.

The pathological features of Wegener's granulomatosis were defined by Godman and Churg (1954). I believe, however, that the renal lesions are simply vascular and

granulomatous lesions as seen in other organs but modified by the local architecture. The basic pathological features of the syndrome are thus: first, giant-cell granulomatous ulceration at one or more levels in the respiratory tract; secondly, widespread giant-cell granulomata, discrete and frequent in lung, kidney, and spleen; thirdly, generalized necrotizing lesions of small vessels, again most common in lung, kidney, and spleen. The initial lesion is that in the respiratory tract, a granulomatous inflammation spreading from the mucosa, but there is little doubt that subsequent thrombosis of small vessels contributes to the development of the large necrotic areas in nares, glottis, and lung. The widespread granulomata resemble those found in experimental serum sickness (Rich, 1942), "allergic granulomatosis" (Churg and Strauss, 1951), and many cases of fatal drug sensitivity (Waugh, 1952; O'Brien and Storey, 1954; Rasmussen, 1955; and others). The necrotizing vascular lesions are morphologically similar to those in polyarteritis nodosa of the "microscopic" variety (Davson et al., 1948), "hypersensitivity angiitis" (Zeek, 1952, 1953), and experimental hypersensitivity (Rich and Gregory, 1943; McKeown, 1947; Crawford and Nassim, 1951; Germuth et al., 1955; and others).

As Fahey et al. (1954) have remarked, Wegener's granulomatosis closely resembles, both clinically and pathologically, several other uncommon diseases "which run in a spectrum from pure necrotizing and granulomatous processes without vasculitis through mixed forms to pure arteritis without granulomas." This close resemblance has led to many cases being described under a wide range of titles (Table I). Klinger (1931), Ringertz (1947), Ahlström et al. (1953), Rose and Spencer (1957), and others regarded their cases as variants of polyarteritis nodosa, whilst many authors have been impressed by the similarity of the local lesion in Wegener's granulomatosis with that in malignant granuloma of the nose (Woodburn and Harris, 1951; Geist and Mullen, 1953; Alexander, 1954; Friedmann, 1955; Paterson, 1956; Singh et al., 1958). Furthermore, Fienberg (1955) was impressed by the similarity of his cases with Loeffler's syndrome and "allergic granulomatosis," and suggested that all three con-ditions be grouped as "pathergic granulomatosis." Yet neither in polyarteritis nodosa, malignant granuloma, Loeffier's syndrome, nor "allergic granulomatosis" is ulceration of the respiratory tract associated with widespread granulomata and vascular necrosis, and I agree with Godman and Churg (1954) in their comment, "The remarkable similarity of the pathological changes in all the cases ... makes it probable that we are dealing with a peculiar and separate syndrome."

Early Course and Development

The early course in Wegener's granulomatosis is one shared by many diseases, ranging from tuberculosis to neo-The diagnosis should be suspected whenever these plasm. can be excluded and destructive lesions in the respiratory tract persist despite antibiotic treatment. When evidence of widespread disease, such as haematuria, uraemia, polyarthritis, and peripheral neuritis, supervenes, the diagnosis is clear. Eosinophilia, conjunctivitis, the characteristic round shadows in chest radiographs, and a haemorrhagic skin rash are confirmatory features. The clinical diagnosis can be supported by biopsies from the ulcerative lesions, which are accessible in the upper air passages in threequarters of the cases, and here eosinophilia, giant cells, and focal granulomata help in the differentiation from malignant granuloma. Biopsy of the skin lesions, which are micro-infarcts due to arteriolar thrombosis, is useful to demonstrate vascular necrosis. In patients in whom the initial or major lesion is in the lung, the obtaining of biopsy material by thoracotomy or bronchoscopy is justifiable in the early stages.

It is clear that two phases, sometimes merging but more often distinct, are involved in the development of Wegener's granulomatosis. The ulceration of the respiratory tract is primary, and the widespread lesions occur later in the natural history of the disease. As polyarteritis nodosa thus cannot be the primary disease, and as no specific microorganism has yet been demonstrated, the aetiology is unknown. Most authors infer that an immunological reaction is involved. Fienberg (1953b) has suggested that the primary lesion is an Arthus phenomenon localized to the bronchial tree, and Kahn (1954, 1955) that the local ulceration is a reaction to an antigen circulating in the blood stream. Yet the evidence is incomplete and the aetiology of the primary lesion must still be regarded as uncertain.

In so far as the widespread, or secondary, lesions are concerned, there is considerable evidence that a hypersensitivity mechanism is involved. First, such a reaction is clearly the causal agent of similar or identical vascular and granulomatous lesions found in diseases such as scarlet fever (Hoyne and Steiner, 1940; Peale et al., 1946) and other infections (Helpern and Trubek, 1933; Contratto, 1947; Bohrod, 1948), asthma (Wilson and Alexander, 1945; Teilum, 1946; Churg and Strauss, 1951), allergic dermatitis (Miale et al., 1947; Rytand et al., 1948), and serum sickness (Clark and Kaplan, 1937; Rich, 1942), in drug reactions (see above), or produced experimentally by measures designed to induce hypersensitivity in animals (see above). Secondly, many of the features of the terminal illness, such as the purpuric skin rash and stomatitis, migratory arthritis, eosinophilia, and transient pareses, are well known to occur in hypersensitivity states (Rich, 1946-7). Thirdly, the raised serum globulin level suggests an antibody response. Finally, impressive evidence came from Case 4, in which symptoms and signs of widespread disease appeared only during and after hypersensitivity reactions to streptomycin (Leggat and Walton, 1956). Widespread lesions morphologically similar to those in the other cases were found at necropsy, but were absent from a lung biopsy made before streptomycin therapy had begun. It is therefore probable that the secondary lesions in Wegener's granulomatosis are due to a hypersensitivity reaction, a drug (Cases 4, 5, 37, and 38) or tissue breakdown products being possible antigens.

Treatment

Many forms of empirical therapy have been unsuccessful. The temporary improvement that sometimes follows antibiotic treatment is probably due to control of secondary infection. Steroids occasionally produce remission (Case 45; Moore *et al.*, 1951; Cutler, 1955), but are more often ineffectual (Cases 39 and 52; Alexander, 1954; Bandler and Campbell, 1954; Paterson, 1956). I believe that the treatment of choice is radiotherapy to the local lesion, with control of secondary infection by suitable antibiotics, and that steroids should if possible be withheld until the local lesion has healed, owing to the danger of hindering the defence against infection of the open respiratory lesions.

Many cases of healing of malignant granuloma of the nose following x rays in low dosage have been reported, and it is logical to try this method on the very similar local lesion in Wegener's granulomatosis. The palatal and nasal lesions in my Cases 9 and 10 were thus healed, and Seidelin and Willcox (1954) and Singh *et al.* (1958) also report success with radiotherapy.

Steroids are the treatment of choice for the widespread lesions once the ulceration of the respiratory tract is healed. They reduce the antibody response (Bjørneboe *et al.*, 1951), inhibit the development of vascular lesions in sensitized animals (Rich *et al.*, 1950; Seifter *et al.*, 1950; Germuth and Ottinger, 1950; and others) and appear to induce healing of the vascular lesions in polyarteritis nodosa (Baggenstoss *et al.*, 1951; Symmers and Litchfield, 1952). The signs of widespread inflammation in Case 9 disappeared following cortisone therapy, the nasal and laryngeal lesions already having been healed by radiotherapy, and the patient has remained well for nearly two years. Cortisone also produced remission of symptoms in Case 10.

Conclusions

The constant clinical course and the presence in all cases of standard anatomical features suggest that the condition under discussion is a separate entity, even though it is closely linked, both morphologically and pathogenetically, with malignant granuloma, periarteritis nodosa, and other granulomatous and vascular diseases. Though the first cases recorded were those of Klinger (1931) and Rössle (1933), it was Wegener (1936, 1939) who first described the disease in detail, and it has come to be known as Wegener's granulomatosis. Including my 10 cases, at least 56 are on record, but many others, less well documented, are probable further examples of the disease.

The natural history clearly has two phases. Beginning with symptoms of chronic nasal or pulmonary infection, it continues with evidence of widespread inflammation. Though a few cases have lived for up to four years, in the majority death rapidly ensues from renal or respiratory failure, the average course being about five months. The clinical diagnosis, though difficult in the early stages, can be confidently made whenever to an ulcerative lesion in the respiratory tract is added evidence of widespread inflammatory disease. The radiological changes in the lungs are typical, and eosinophilia and biopsy of the local lesion or of skin are of confirmatory value.

The diagnostic pathological features are granulomatous ulceration at any level in the respiratory tract and widespread giant-cell granulomata and necrotizing lesions of small vessels. The characteristic round or oval consolidations in the lungs are not infarcts but conglomerate areas of peribronchial necrosis. The severe and constant changes in the renal corpuscles are focal, do not represent diffuse glomerulonephritis, but are simply granulomatous and vascular lesions modified by the local architecture.

The initial lesion in Wegener's granulomatosis is not polyarteritis nodosa, but the ulceration in the respiratory tract, the actiology of which is uncertain. The widespread lesions occur later in the course of the disease and there is impressive evidence that they are the result of a hypersensitivity reaction. Treatment should be primarily directed towards healing of the initial lesion, and radiotherapy has proved successful in some cases. Administration of cortisone or other steroids is the treatment of choice once widespread lesions have occurred.

Summary

Wegener's granulomatosis is an uncommon syndrome in which giant-cell granulomata in the respiratory tract occur together with granulomatous and vascular lesions resembling those in polyarteritis nodosa. In this work, which is based on a study of 10 cases, and 46 others selected from the literature, the clinical and pathological features are described and tabulated.

The disease is clearly a separate entity, differing both clinically and in the character and distribution of the lesions from similar conditions, such as polyarteritis nodosa and malignant granuloma. It begins as a progressive ulceration of unknown aetiology in the respiratory tract. Sooner or later widespread lesions, the result of a hypersensitivity reaction, complete the disease picture. Treatment of the lesions of the respiratory tract by radiotherapy has been successful in a few cases.

ADDENDUM.-Since this paper was written, I have seen further reports of cases of Wegener's granulomatosis by Fanger and Hoffman (1957), Plummer et al. (1957), Levine and Madden (1957), Read and Treip (1957), Stoeckle et al. (1957), and Gordon et al. (1957).

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