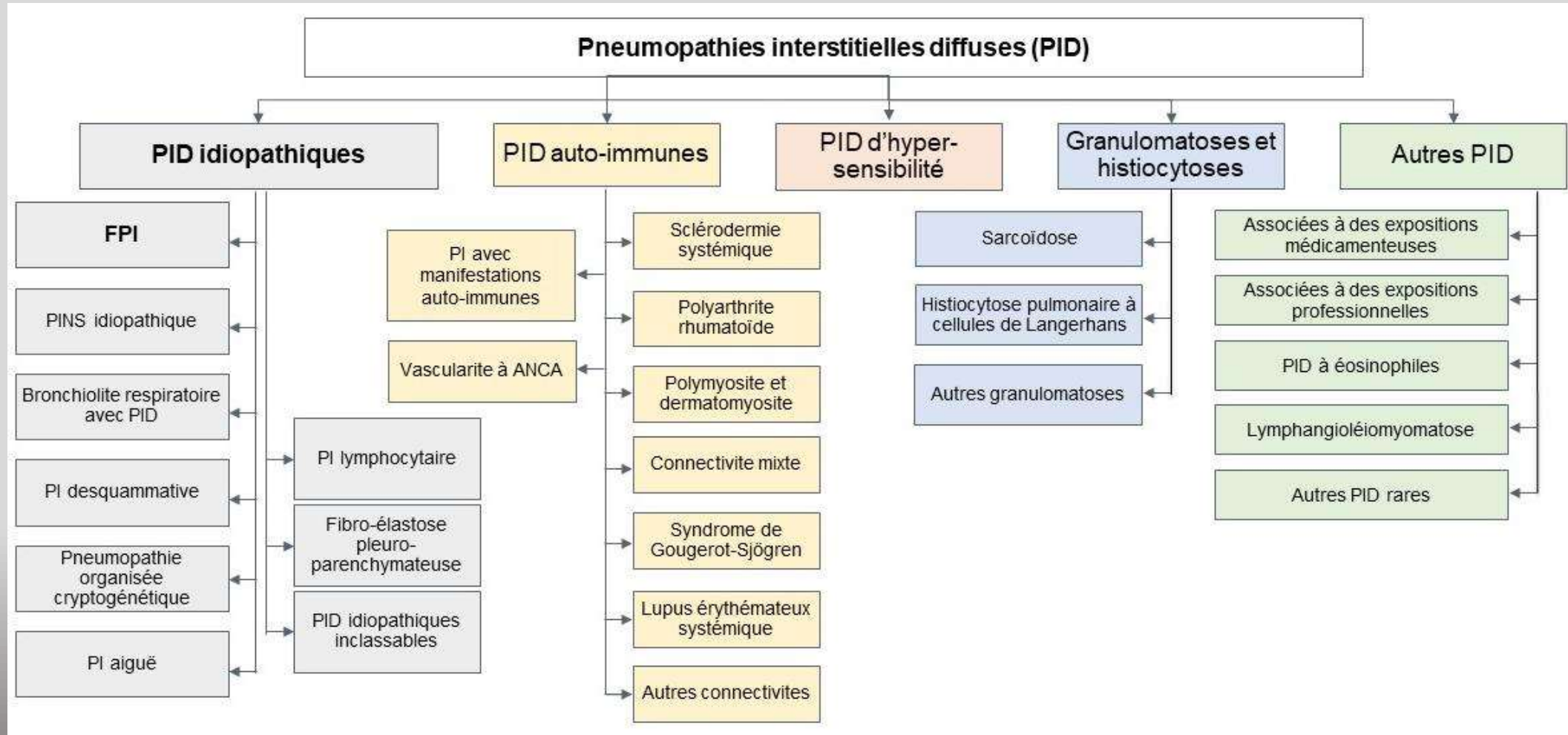


Pneumopathie d'hypersensibilité chronique

Catherine Beigelman CHUV Lausanne
Guillaume Prévot CHU Toulouse

Classification usuelle des PID



Définition de la PHS

- Maladie inflammatoire et/ou fibreuse affectant le parenchyme pulmonaire et les petites VA
- Typiquement : réaction immuno-médiée après exposition à un antigène inhalé
- Antigène non identifié dans 30 à 50% des cas ...

- Aiguë / Subaiguë / Chronique : termes abandonnés

PHS fibreuse

Ou

PHS non fibreuse

Epidémiologie des PHS

Table 2. ILD Diagnosis among Patients in ILD-India Registry Cohort

Type of ILD	Number of Patients (%)
Hypersensitivity pneumonitis	513 (47.3)
Connective tissue-associated ILD	151 (13.9)
Idiopathic pulmonary fibrosis	148 (13.7)
Idiopathic nonspecific interstitial pneumonia	92 (8.5)
Sarcoidosis	85 (7.8)
Pneumoconiosis	33 (3.0)

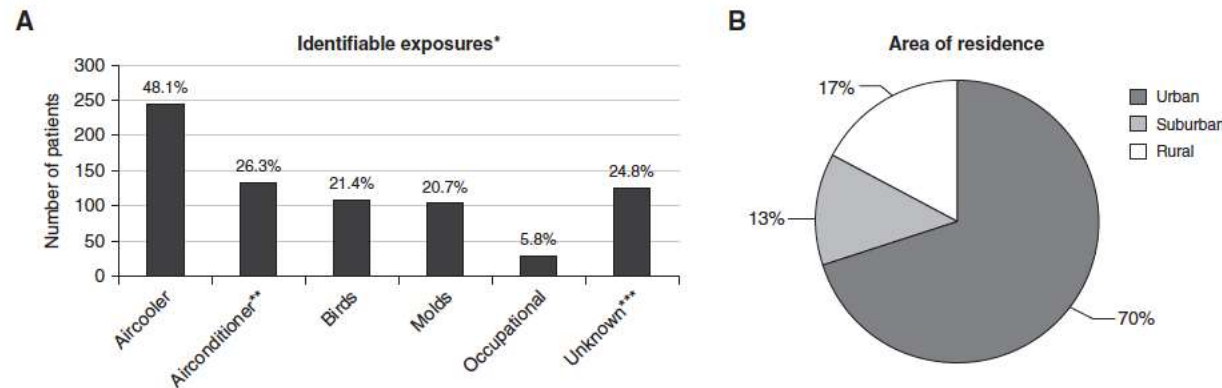
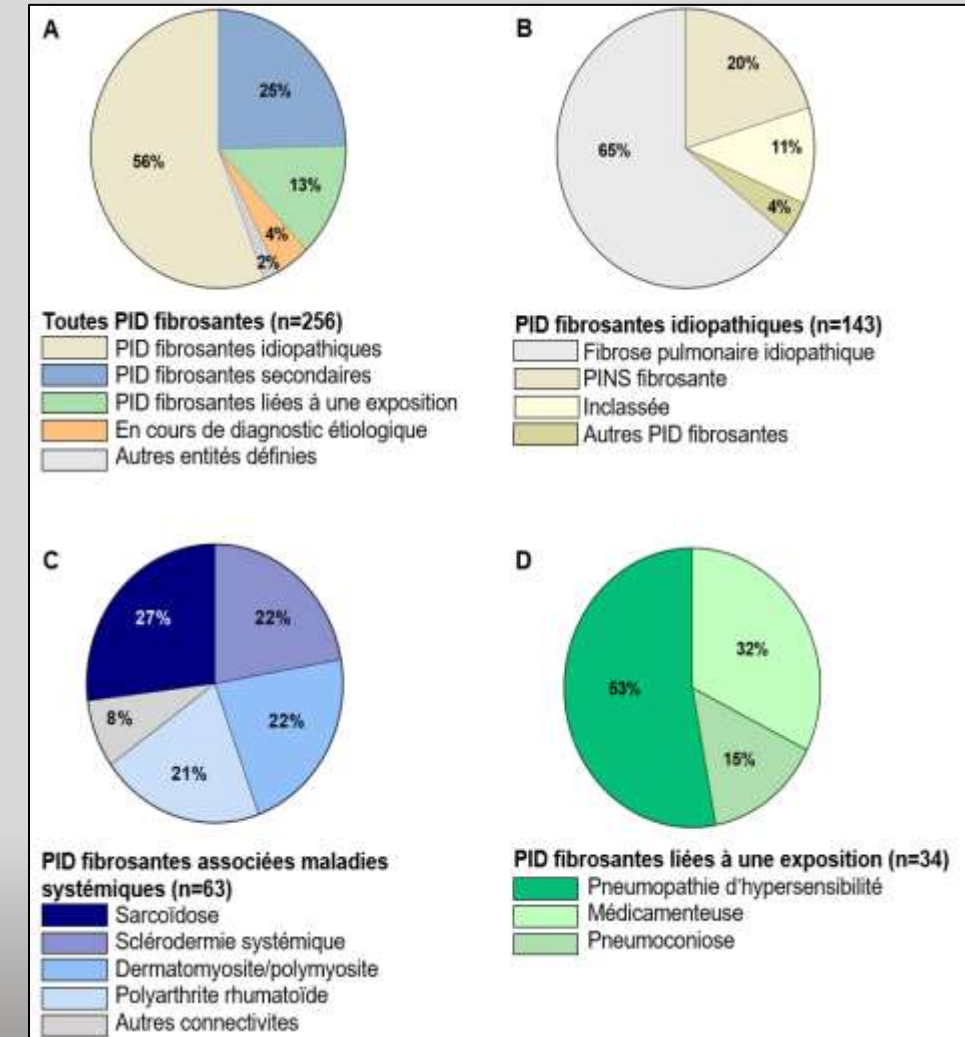
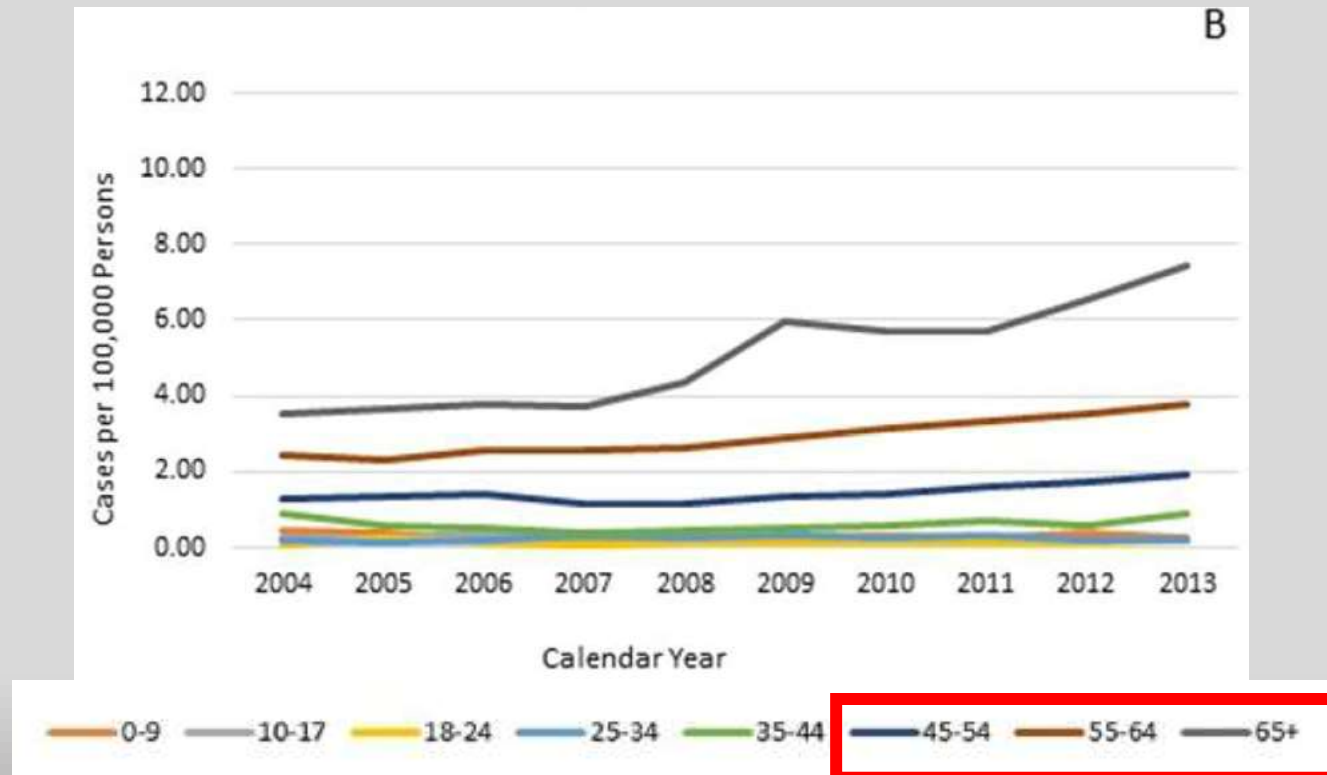


Figure 5. (A) Identifiable exposures among 513 patients with hypersensitivity pneumonitis (HP). (B) Patients with HP (n = 513) by area of residence. *Numbers do not add to 100% because some patients had multiple exposures. **Although it is unknown if exposure to air conditioners is associated with HP, general practice in India is not to clean air conditioners on a regular basis, so it is conceivable that mold could grow in these units. People with air conditioners may then have occult exposure to mold dispersed into the environment. ***24.8% of patients with HP did not have overt exposures to known causes of HP, which is in keeping with reports of insidious or unknown exposures in the literature (42, 43).

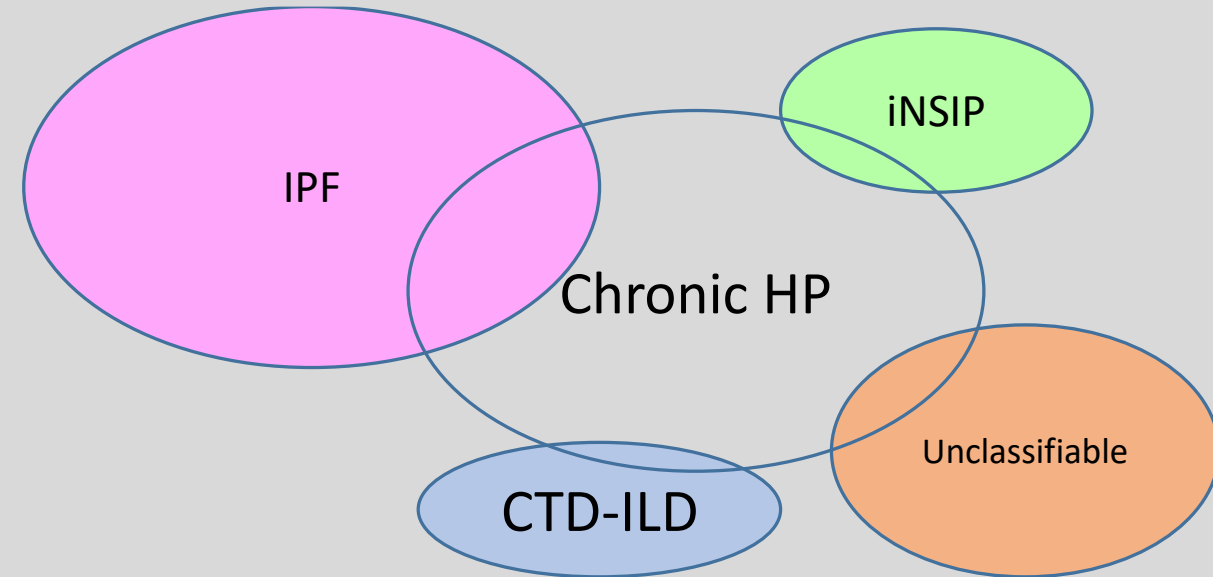


Prevalence of chronic hypersensitivity pneumonitis



Prévalence estimée à
2/100000 hbt aux US

PHSf : un diagnostic difficile



Chronic hypersensitivity pneumonitis in patients diagnosed with idiopathic pulmonary fibrosis: a prospective case-cohort study

Ferran Morell, Ana Villar, María-Angela Montero, Xavier Muñoz, Thomas V Colby, Sudhakar Pipvath, María-Jesús Cruz, Ganesh Raghu

Lancet Respir Med 2013

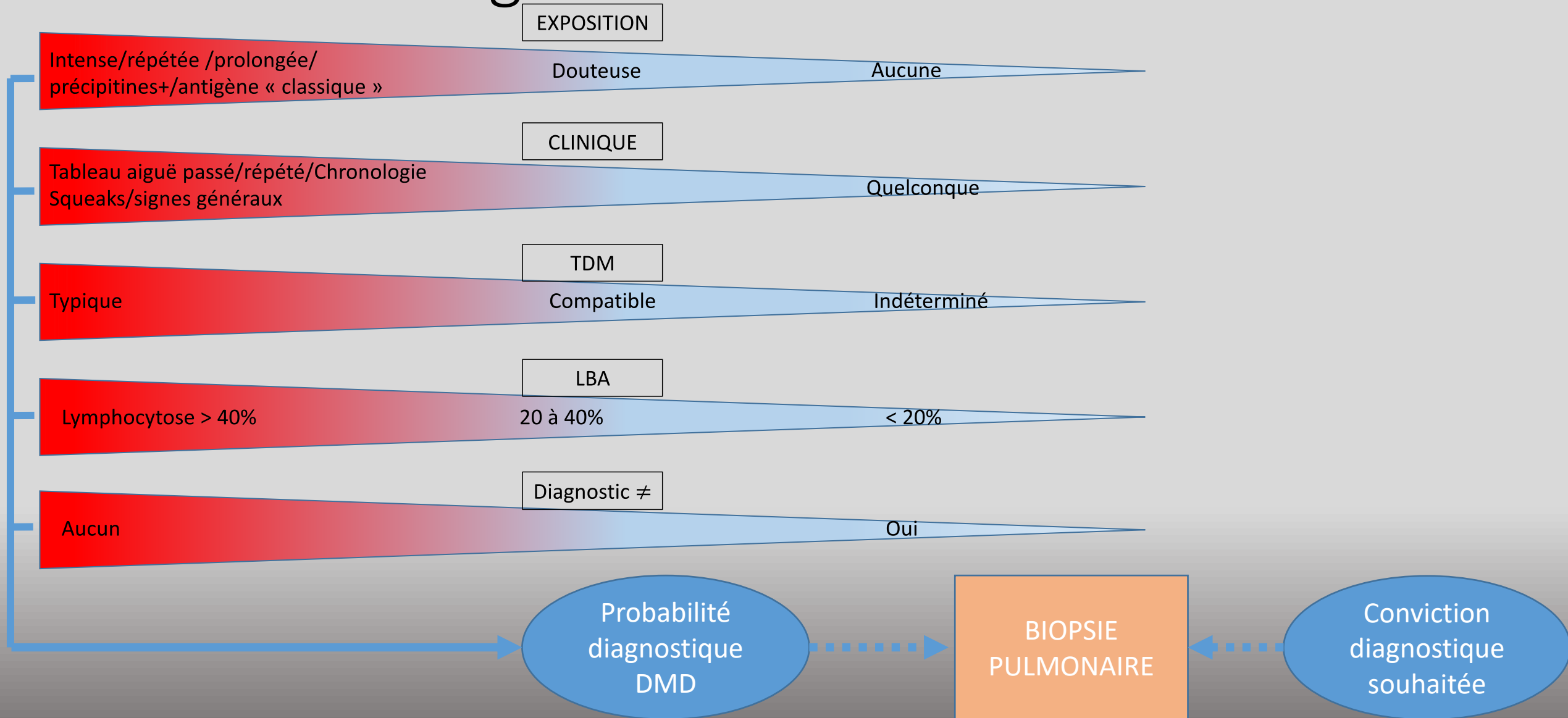
	Clinicians (κw)	Radiologists (κw)	Pathologists (κw)	MDTM (κw)
Idiopathic pulmonary fibrosis	0.72 (0.67–0.76)	0.60 (0.46–0.66)	0.58 (0.45–0.66)	0.71 (0.64–0.77)
Connective tissue disease-related interstitial lung disease	0.76 (0.70–0.78)	0.17 (0.08–0.31)	0.21 (0.06–0.36)	0.73 (0.68–0.78)
Non-specific interstitial pneumonia	0.31 (0.27–0.41)	0.32 (0.26–0.41)	0.30 (0.00–0.53)	0.42 (0.37–0.49)
Hypersensitivity pneumonitis	0.42 (0.30–0.47)	0.35 (0.29–0.43)	0.26 (0.10–0.45)	0.29 (0.24–0.40)

Data are median (IQR). MDTM=multidisciplinary team meeting.

Table 4: Weighted kappa values (κw) for estimation of diagnostic likelihood for individual diagnoses of diffuse parenchymal lung disease

Lancet Respir Med 2016

PHSf : un diagnostic difficile



Recherche de la source antigénique

TABLE 3 Evidence-based exposure screening questionnaire

			Number of published cases	Number of publications
1. Have you been exposed to birds or feather/down-containing items?				
☐ YES	☐ NO	Bird: Pet birds (tropical), pigeon breeding or other birds in your environment? (duck and geese)	243	26
☐ YES	☐ NO	Feather/down-containing items (duck or goose down-containing jackets, pillows, blankets or feather dusters)?	111	7
☐ YES	☐ NO	Exposure to bird droppings in a farm, factory, or home setting (droppings in an attic, barn, porch or yard)?	2	1
2. Have you had any of the following in your home or work environment?				
☐ YES	☐ NO	Humidifiers or mist fountains	17	5
☐ YES	☐ NO	Mould or mildew	13	10
☐ YES	☐ NO	Moist or decayed wood	9	6
☐ YES	☐ NO	Flood/water damage	4	4
☐ YES	☐ NO	Straw mats	2	1
☐ YES	☐ NO	Window or single unit air conditioners	1	1
☐ YES	☐ NO	Oil fan heater	1	1
☐ YES	☐ NO	Steam iron	1	1
3. Do you frequently use a hot tub, jacuzzi or sauna?				
☐ YES	☐ NO		58	17
4. Have you had any of the following occupations or hobbies or worked in any of the following locations?				
☐ YES	☐ NO	Farm/greenhouse worker	295	20
☐ YES	☐ NO	Machine operator	48	5
☐ YES	☐ NO	Mushroom worker or worker in mushroom factories	42	8
☐ YES	☐ NO	Carpenter/sawmill worker or worked in a hardwood processing plant	16	5
☐ YES	☐ NO	Wind or brass musical instrument (e.g. saxophone, bassoon, tenor horn, bagpipe)	5	4
☐ YES	☐ NO	Painter or paint sprayer	5	2
☐ YES	☐ NO	Garbage collector	5	2
☐ YES	☐ NO	Well digger	5	1
☐ YES	☐ NO	Baker	3	2
☐ YES	☐ NO	Working with plaster	1	1
☐ YES	☐ NO	Plastic welder	1	1
☐ YES	☐ NO	Food production or processing: salami, dry sausage, green tea, onion, potato, flour, wheat, seafood	17	9
☐ YES	☐ NO	Cork factory	17	2
☐ YES	☐ NO	Repair garage or aircraft manufacturing	10	4
☐ YES	☐ NO	Esparto fibre factory	10	2
☐ YES	☐ NO	Cosmetic production	2	1
☐ YES	☐ NO	Warehouses	1	1
☐ YES	☐ NO	Feeding stores/factory	1	1

Mold in Foam Pillows and Mattresses A Novel Cause of Hypersensitivity Pneumonitis

Check for updates

Onofre Moran-Mendoza, MD, PhD; Sharina Aldhaheri, MD; Connor J. A. Black, BSc (Hons); Marie Clements-Baker, MD; Mohamed Khalil, MD; and Alexander Boag, MD

Hypersensitivity pneumonitis (HP) is an inflammatory and/or fibrotic disease affecting the lung parenchyma and small airways. It typically results from an immune-mediated reaction provoked by an overt or occult inhaled antigen in susceptible individuals. The chronic or fibrotic form of HP has a poor prognosis, especially when no inciting antigen is identified, which occurs in up to 60% of cases. We report two cases of HP associated with exposure to mold in foam pillows and a mattress, which has not previously been reported as a risk factor for HP. Given the high prevalence of foam in pillows and mattresses, mold in foam in bedding may explain many HP cases with a previously unrecognized cause. Early identification and avoidance of foam in bedding may prevent HP progression to end-stage pulmonary fibrosis and death.

CHEST 2021; 160(3):e259-e263

PO17-428

Service de pneumologie
Pour des soins personnalisés

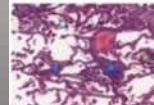
Une pneumonie d'hypersensibilité par ingestat
V Dongay, L Mhanna, G Prévot, E Noël-Savina, M Colombat, L Guillenninault, S Pontier – Marchandise
Toulouse University Hospital - Toulouse (France)

TDM d'entrée au 2^{ème} épisode
23/04/20

Eviction, antibiothérapie, diurétiques
04/05/20

Le diagnostic de pneumopathie d'hypersensibilité suite à l'ingestion de *Scutellaria Batcalensis* a été retenu devant :

- La biopsie pulmonaire chirurgicale qui suggère un phénomène immuno-allergique sans atteinte bronchiale.
- L'absence d'argument pour une participation infectieuse ou cardiogénique.
- Le TDM thoracique retrouvant un aspect non spécifique mais compatible avec une PHS
- Absence de rechute suite à l'arrêt des compléments nutritionnels malgré le retour à domicile



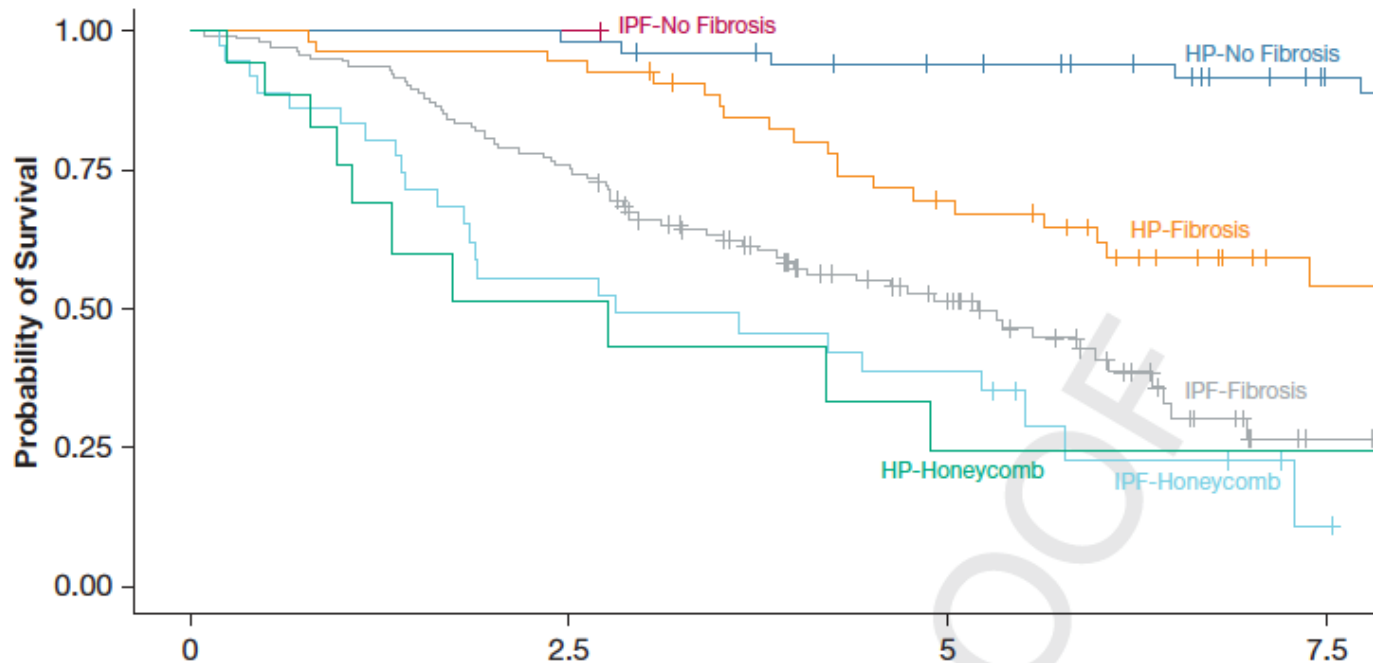


Pronostic de la PHSf

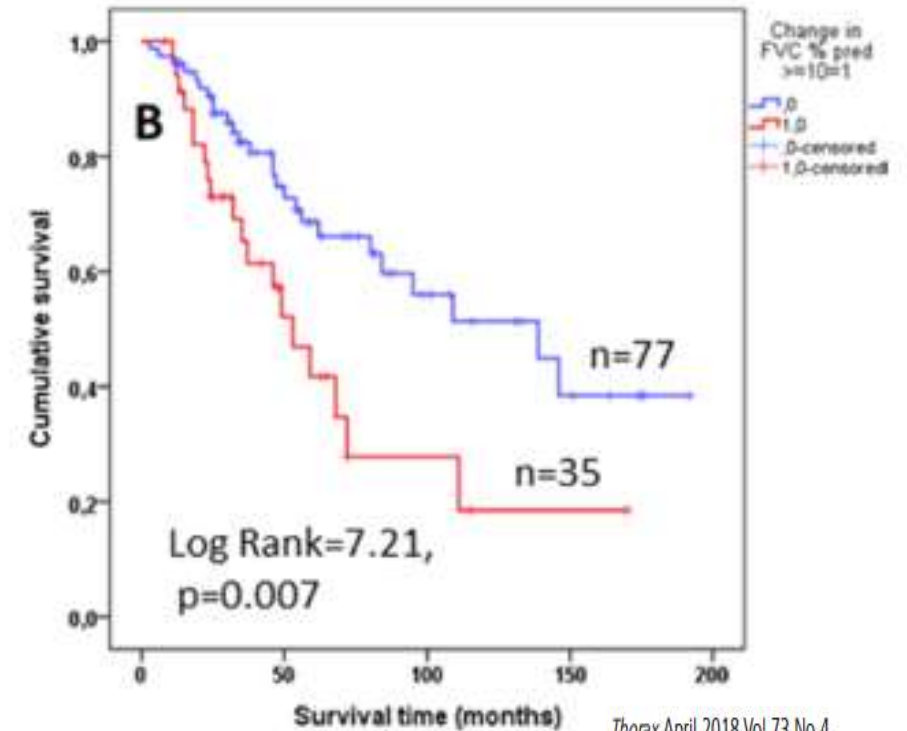
Hypersensitivity Pneumonitis

Radiologic Phenotypes Are Associated With Distinct Survival Time and Pulmonary Function Trajectory

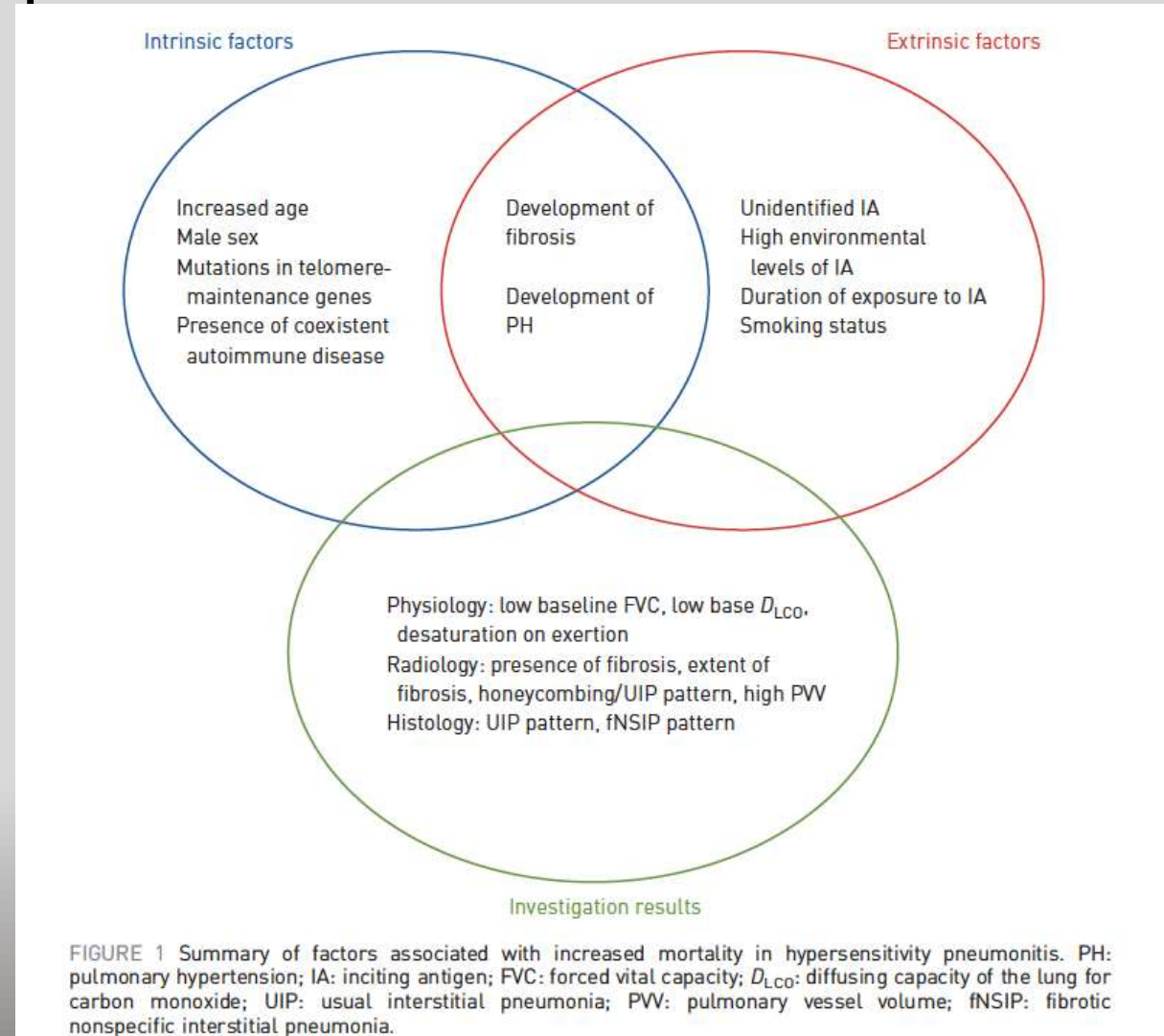
Chest 2019



B). Survival time estimate of chronic hypersensitivity pneumonitis cohort based on decrease $<10\%$ ($n=77$) and $\geq 10\%$ predicted FVC ($n=35$) after 6–12-month period of follow-up



Facteurs pronostics des PHS



Généralités

Non fibrotique

Phénomènes inflammatoires et souvent réversibles

Fibrotique

PHS chronique + impact diagnostique de certains signes RX



Formes aiguës, subaiguës,
chroniques

Rappels

ATTENUATION EN MOSAÏQUE

Hétérogénéité des densités pulmonaires en inspiration

Maladie vasculaire

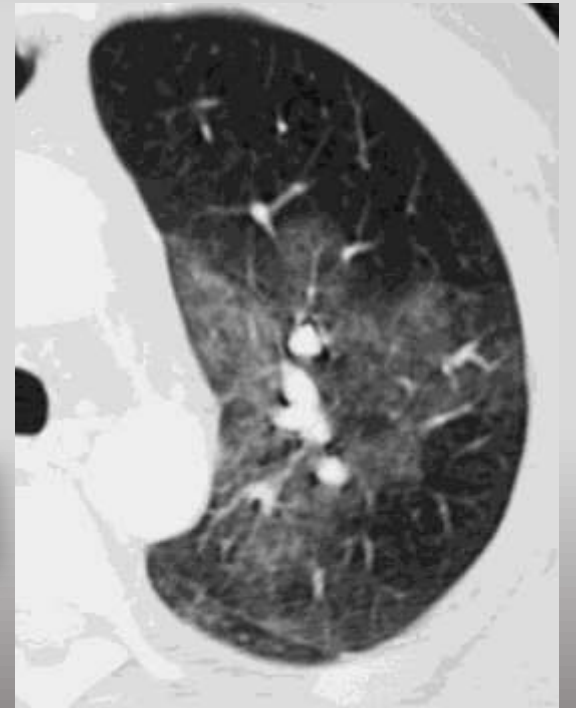
Atteinte des voies aériennes

Infiltration interstitielle ou alvéolaire en verre dépoli



2 situations principales
Hyper-Hypo atténuation

Taille des vaisseaux pulmonaires ++



Approche diagnostique

Les vaisseaux ont-ils un calibre équivalent dans les zones hypo et hyperdenses?



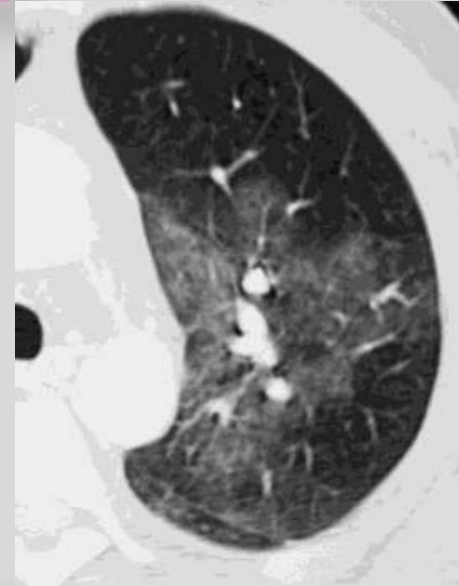
Oui

Pathologie infiltrante



Non

**Pathologie vasculaire
Pathologie des PVA**



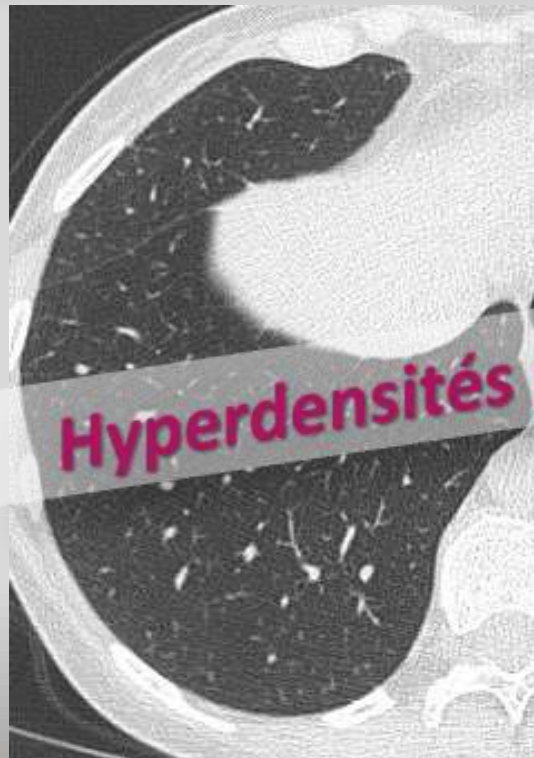
Verre dépoli en mosaïque

Vascularisation pulmonaire uniforme

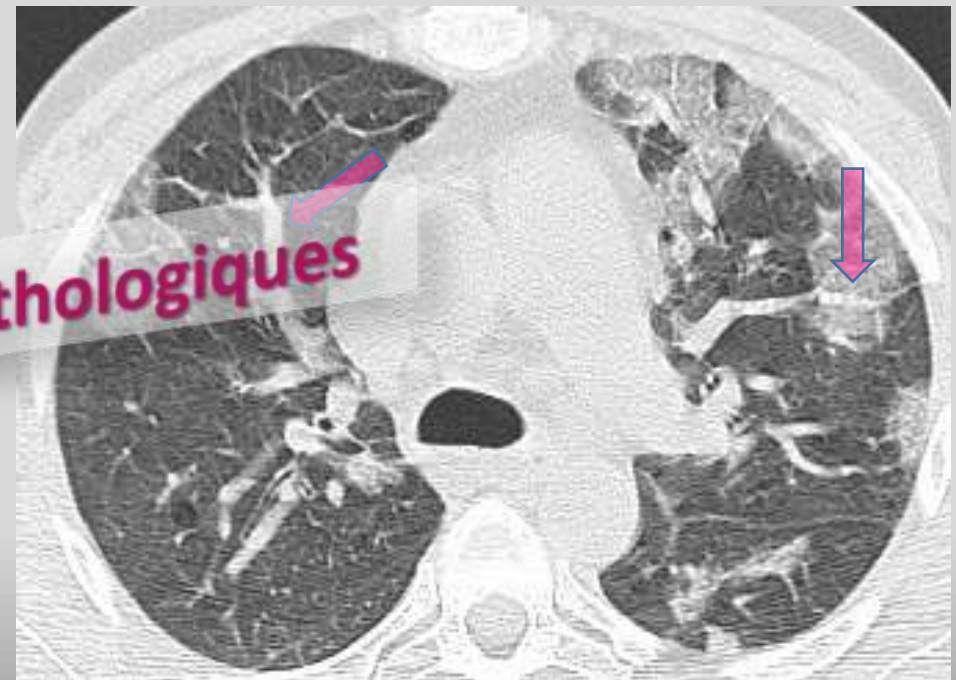
Pathologie infiltrante aiguë, subaiguë ou chronique



PINS



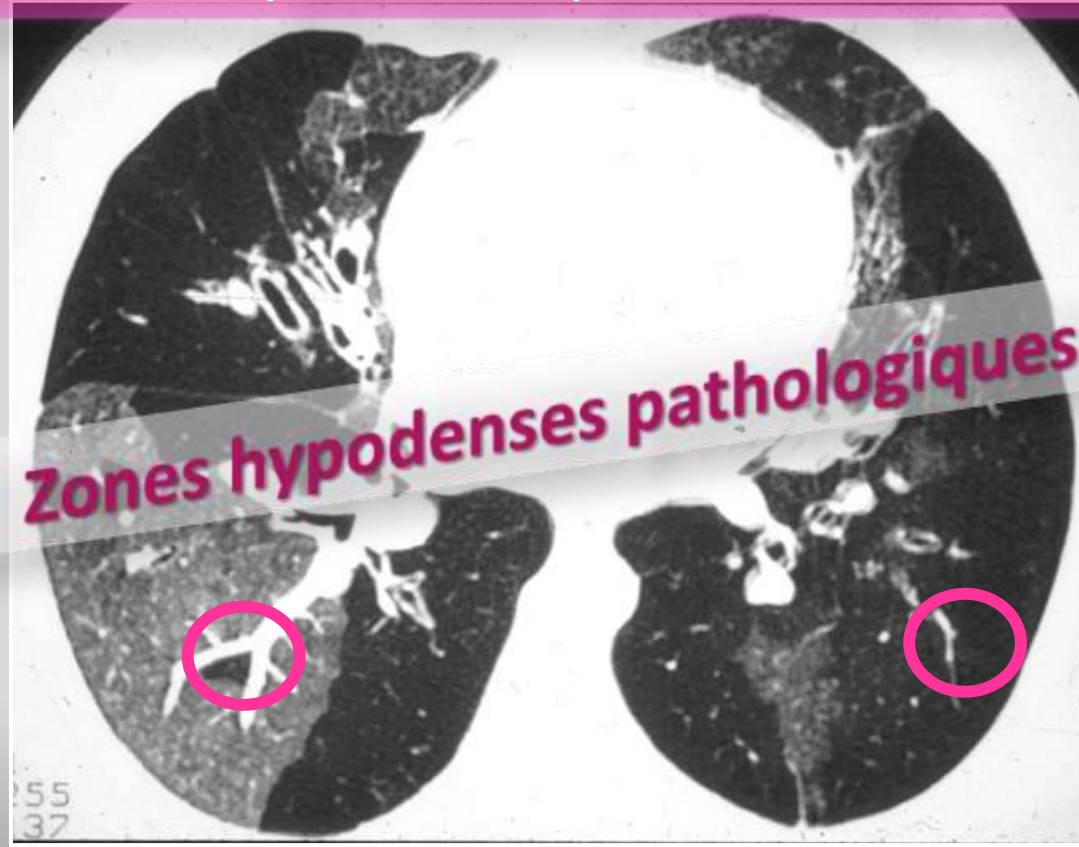
Hyperdensités pathologiques



Infection à Staphylocoque et Haemophilus

Perfusion en mosaïque

Différences régionales de densité secondaires à des différences régionales de perfusion pulmonaire



Remerciement P.Grenier

Zones en verre dépoli

Hypoperfusion
↘ taille vx pulmonaires

↗ flux sg capillaire par
redistribution vasculaire

Perfusion en mosaïque

Cause bronchiolaire

Bronchiolite constrictive



Cause vasculaire primitive

CPC post-embolique

Anomalies des voies aériennes coassociées

Dilatation AP/ Cavités droites

Expiration ?

Piégeage

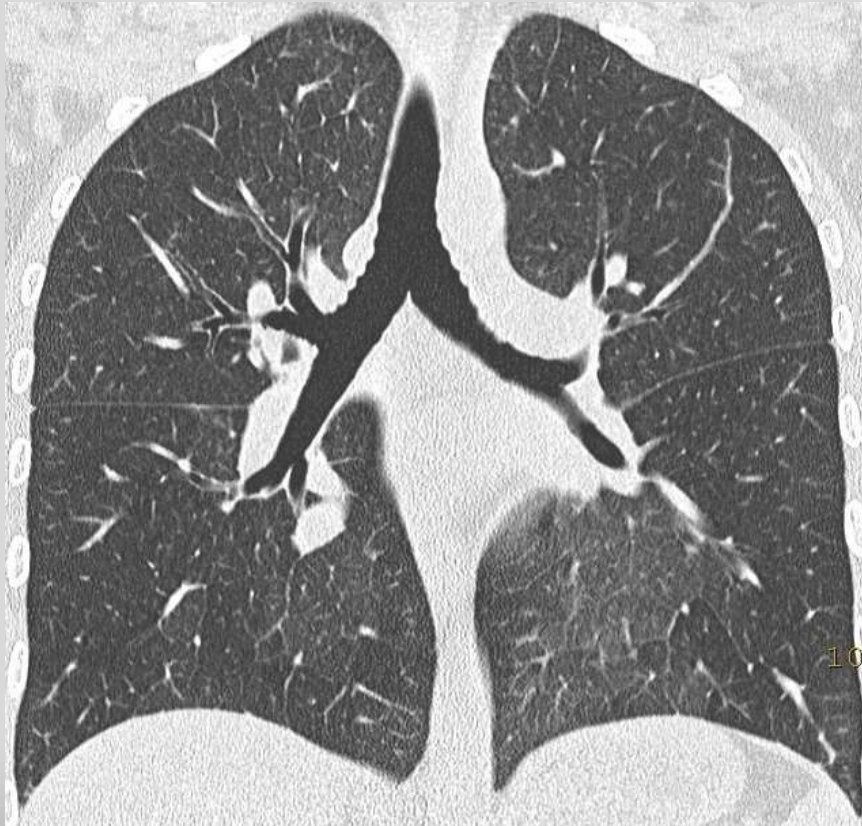
Pas de Piégeage

Même gradient d'atténuation entre hypo/hyper

↗ différences atténuation

Piégeage expiratoire

Zones focales d'hypoatténuation au sein du poumon normal en hyperdensité en expiration



Bronchiolite constrictive post-allogreffe de moëlle

Aspect en mosaïque des densités pulmonaires

Verre dépoli en mosaïque

Pneumonie interstitielle

Lignes septales
Réticulations
Bronchectasies
Images kystiques

Perfusion en mosaïque

M. des voies aériennes

Piégeage
Bronchectasies

M. vasculaires

Dilatation des artères
pulmonaires

Aspect en fromage de tête

Pneumonie interstitielle et maladie des voies aériennes

Aspect en fromage de tête

Pneumonie interstitielle + Maladies des voies aériennes

Pneumonie d'hypersensibilité

Sarcoïdose

BR-PI/DIP

Bronchiolite folliculaire/LIP

Infections atypiques

OAP + asthme



Non spécifique de PHS non fibrosante
Spécifique de PHS fibrosante

PHS fibrosante typique

Table 6. Chest HRCT Scan Features of the Fibrotic HP Pattern

HRCT Pattern	Typical HP	Compatible with HP	Indeterminate for HP
Description	The “typical HP” pattern is suggestive of a diagnosis of HP. It requires a) an HRCT pattern of lung fibrosis (as listed below) in one of the distributions and b) at least one abnormality that is indicative of small airway disease	“Compatible-with-HP” patterns exist when the HRCT pattern and/or distribution of lung fibrosis varies from that of the typical HP pattern; the variant fibrosis should be accompanied by signs of small airway disease	The “indeterminate-for-HP” pattern exists when the HRCT is neither suggestive nor compatible with a typical and probable HP pattern
Relevant radiological findings	HRCT abnormalities indicative of lung fibrosis are most commonly composed of irregular linear opacities/coarse reticulation with lung distortion; traction bronchiectasis and honeycombing may be present but do not predominate The distribution of fibrosis may be: • Random both axially and craniocaudally or • Mid lung zone–predominant or • Relatively spared in the lower lung zones HRCT abnormalities indicative of small airway disease: • Ill-defined, centrilobular nodules and/or GGOs • Mosaic attenuation, three-density pattern,* and/or air trapping (often in a lobular distribution)	Variant patterns of lung fibrosis: • UIP pattern: basal and subpleural distribution of honeycombing with/without traction bronchiectasis (per 2018 diagnosis of IPF guidelines [20]) • Extensive GGOs with superimposed subtle features of lung fibrosis Variant (predominant) distributions of lung fibrosis: • Axial: peribronchovascular, subpleural areas • Craniocaudal: upper lung zones HRCT abnormalities indicative of small airway disease: • Ill-defined centrilobular nodules, or • Three-density pattern* and/or air trapping	Lone patterns (i.e., not accompanied by other findings suggestive of HP) of: • UIP pattern (as per 2018 IPF diagnosis guidelines [20]) • Probable UIP pattern (as per 2018 IPF diagnosis guidelines [20]) • Indeterminate pattern for UIP (as per 2018 IPF diagnosis guidelines [20]) • Fibrotic NSIP pattern • Organizing pneumonia–like pattern • Truly indeterminate HRCT pattern

Definition of abbreviations: GGO = ground-glass opacity; HP = hypersensitivity pneumonitis; HRCT = high-resolution computed tomography; IPF = idiopathic pulmonary fibrosis; NSIP = nonspecific interstitial pneumonia; UIP = usual interstitial pneumonia.

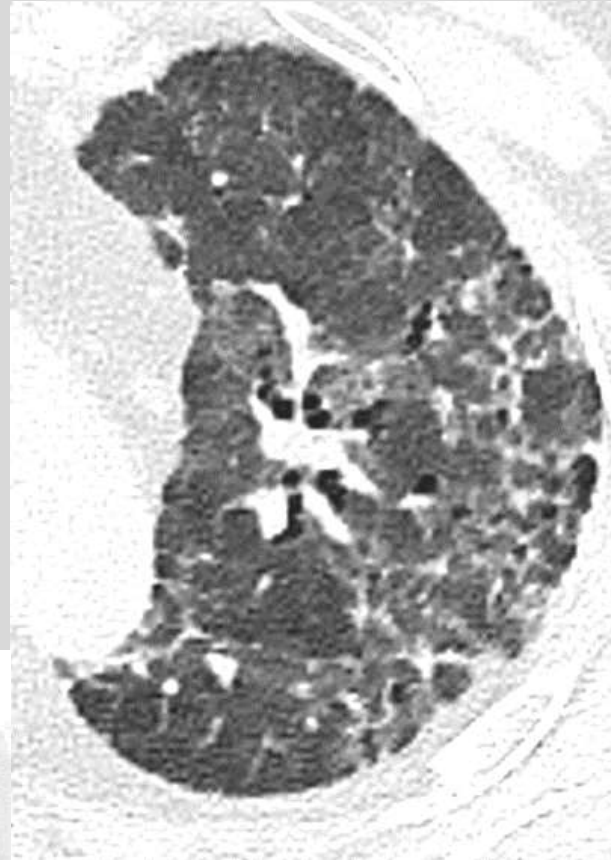
Rarely, fibrotic HP may be seen 1) as a component of combined pulmonary fibrosis and emphysema or pleuroparenchymal fibroelastosis with emphysema, 2) as a pure emphysematous form of HP, or 3) in acute exacerbation.

*The three-density pattern was formerly called the “headcheese sign.” It is described in detail in Table 4.

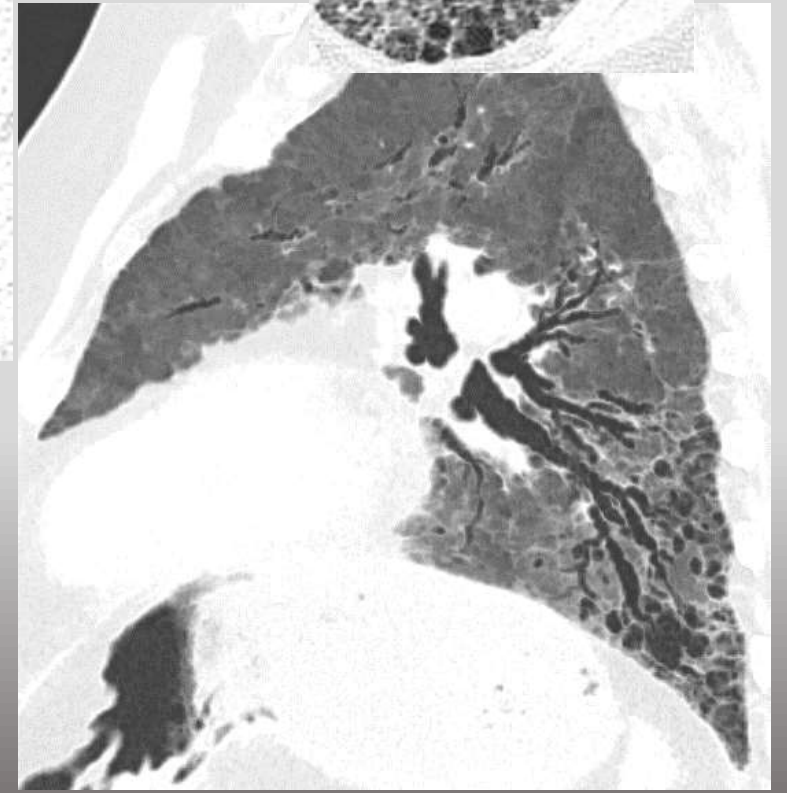
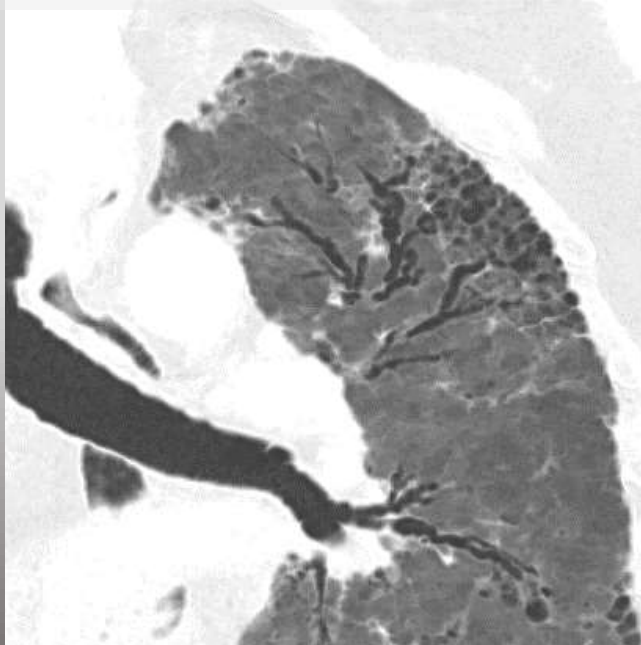
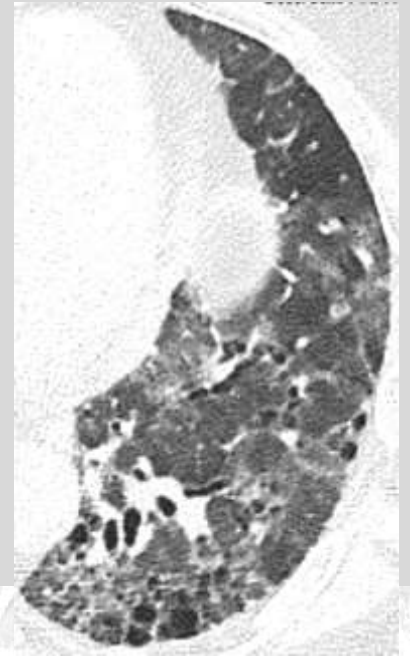
Fibrose



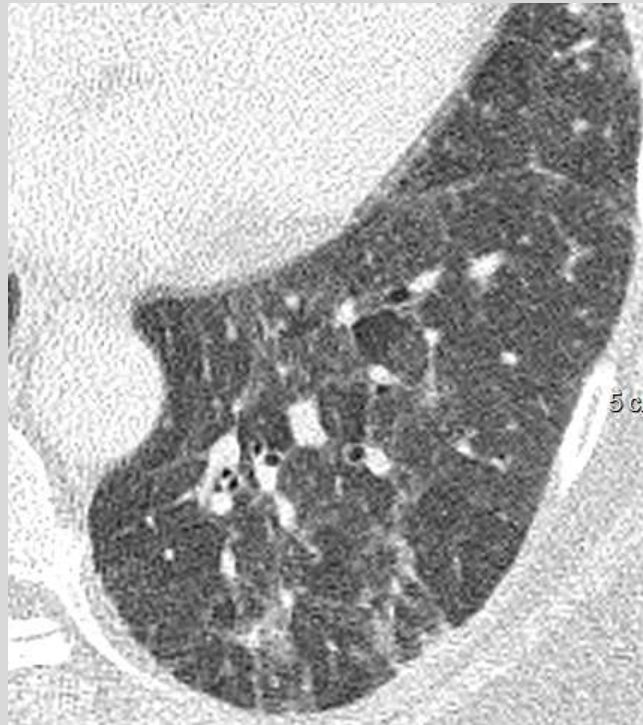
Réticulations irrégulières
Distorsion architecturale
Bronchectasies traction



Sans
prédominance
crânio-caudale

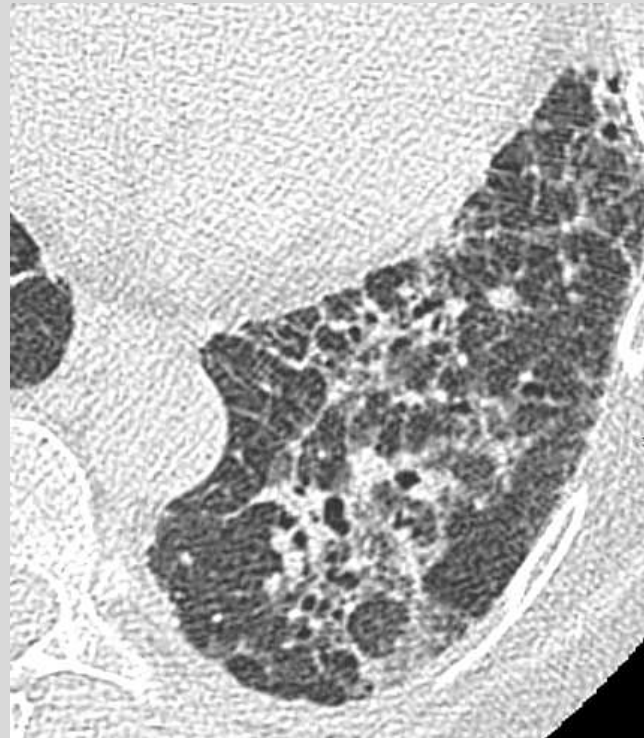


18/01/13



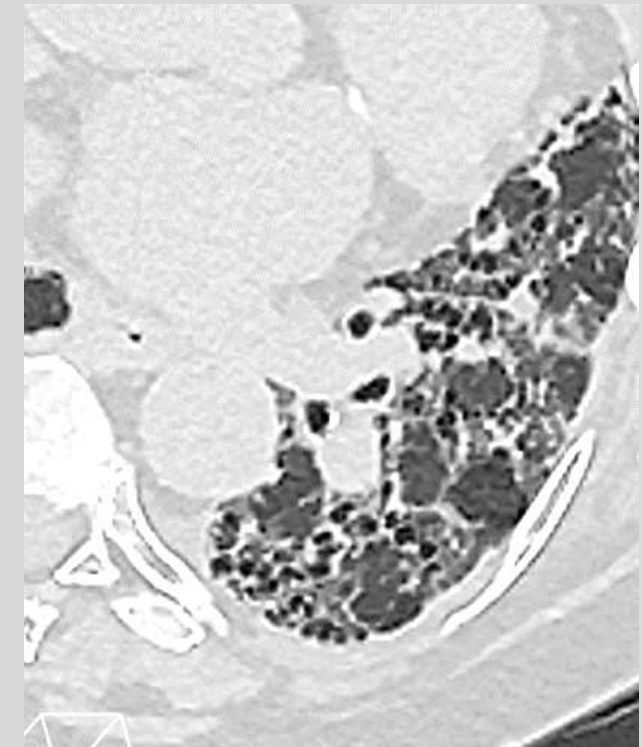
Verre dépoli
Hypoatténuation

05/05/17

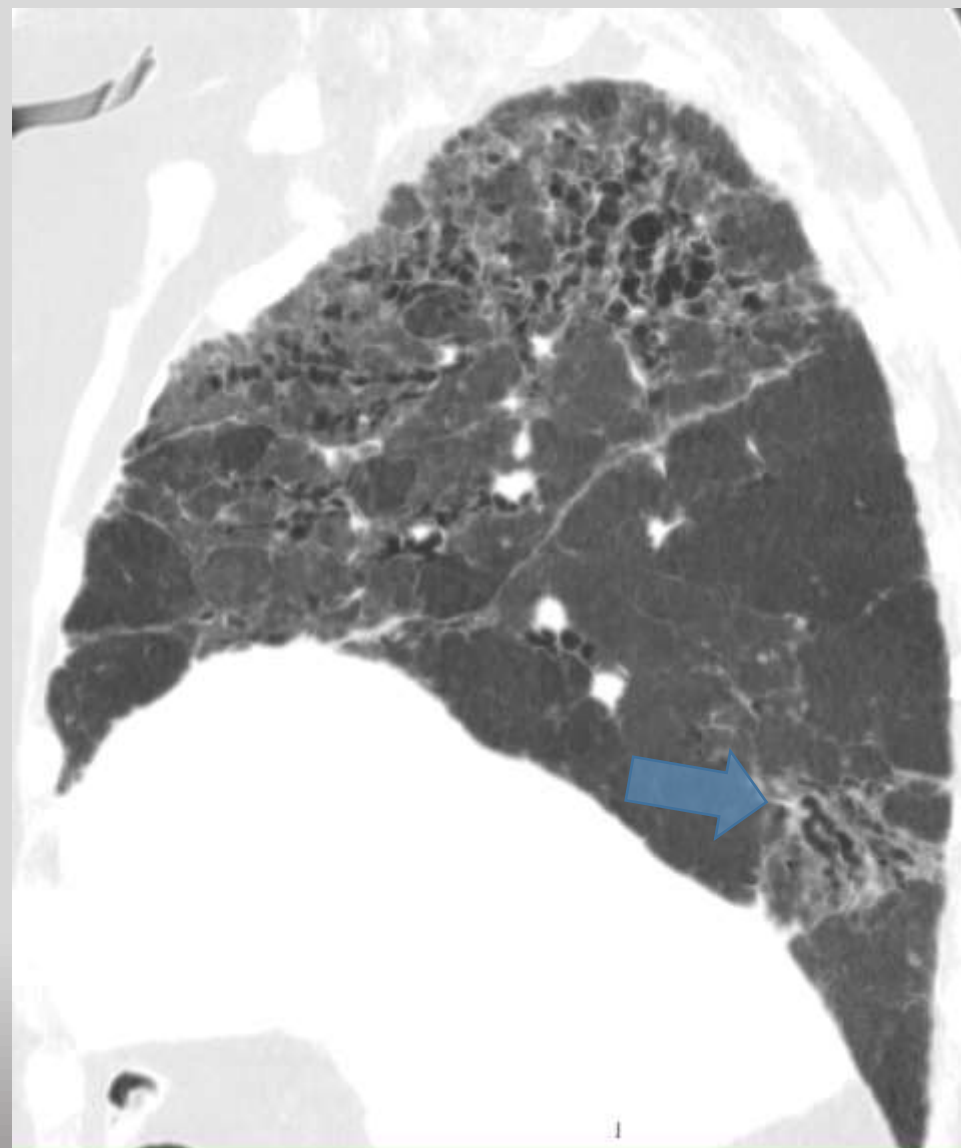
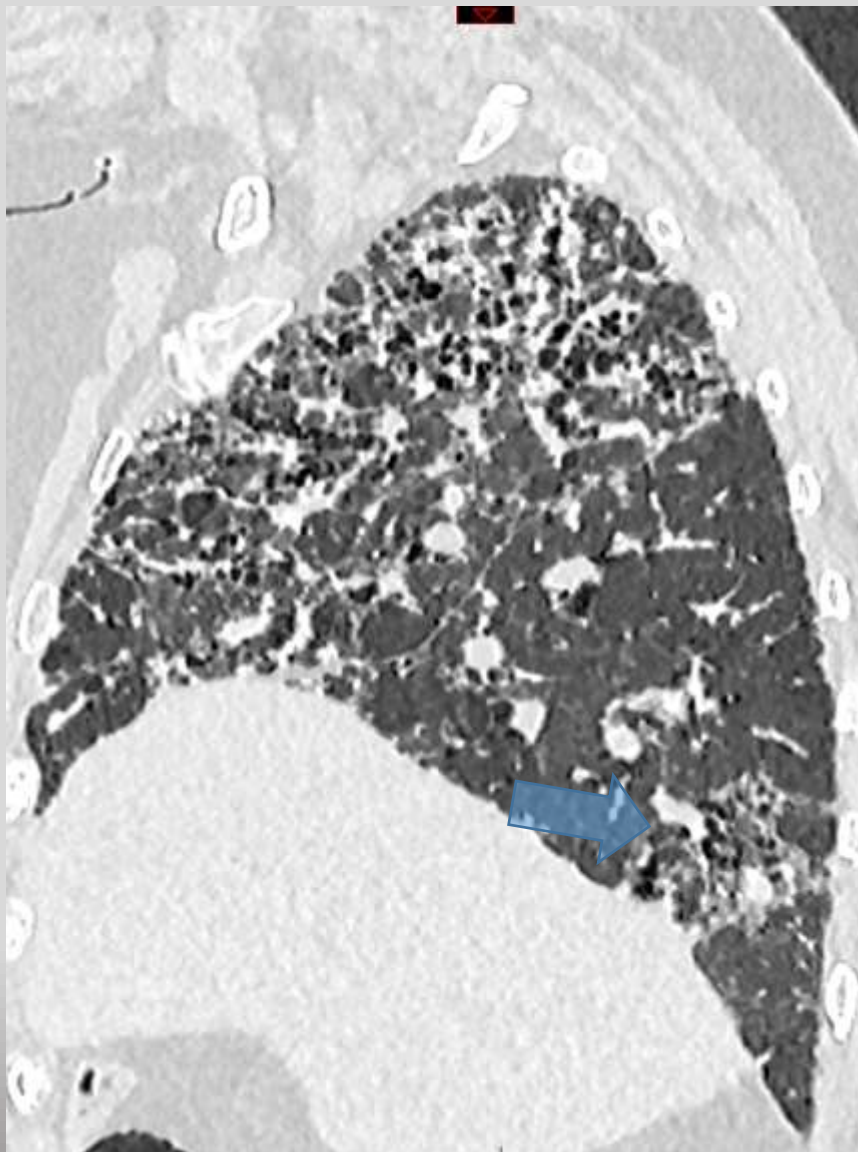


Réticulations irrégulières
Distorsion architecturale
Hyperdensités avec DDB

18/06/21



Rayon de miel
Perte de volume



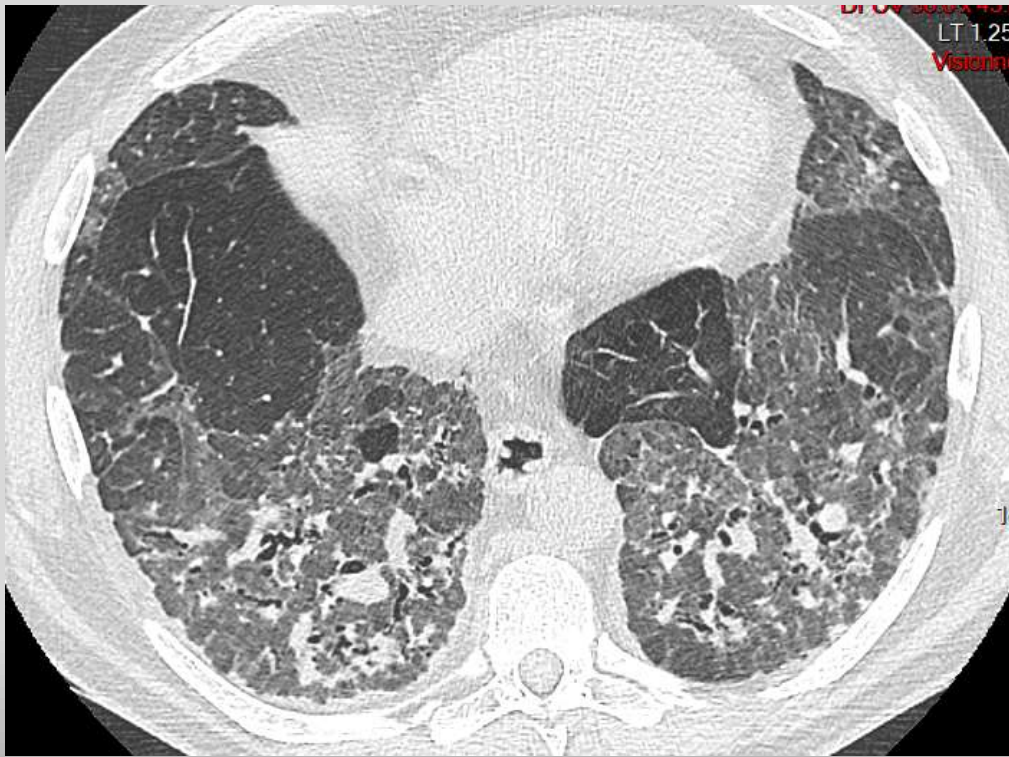
Moindre atteinte des zones inférieures

Atteinte des petites voies aériennes

Atténuation en mosaïque de forme lobulaire: perfusion en mosaïque

Three-density pattern

Et/ou piégeage expiratoire



1mm Lung

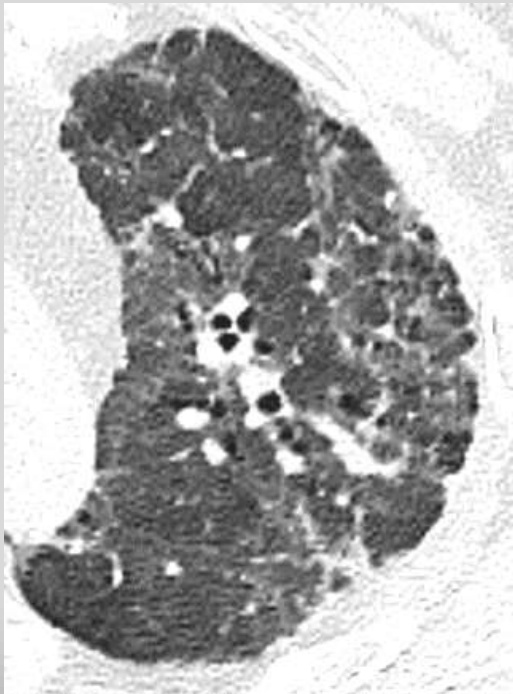


Expiration

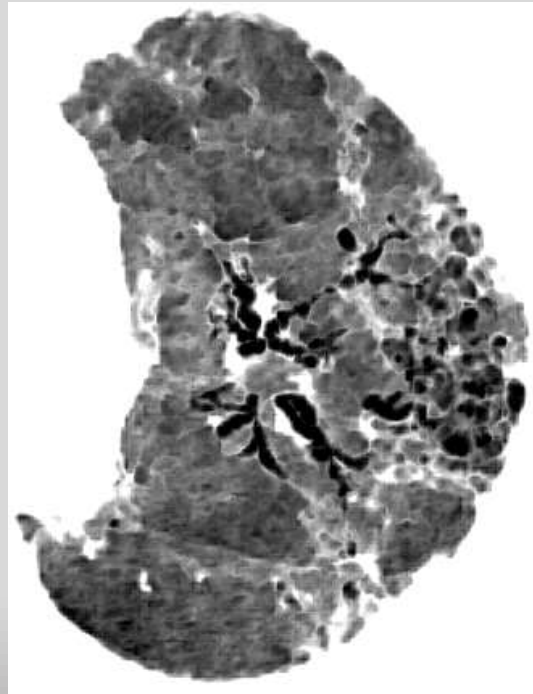
Poumon des boulangers

**Fibrose
SAD?**

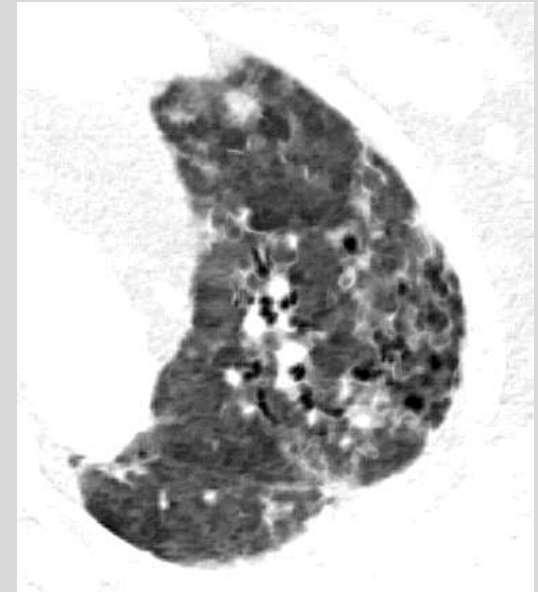
**Sensibilisation des différences de
densité en filtre mou et mIP**



Lung 1 mm

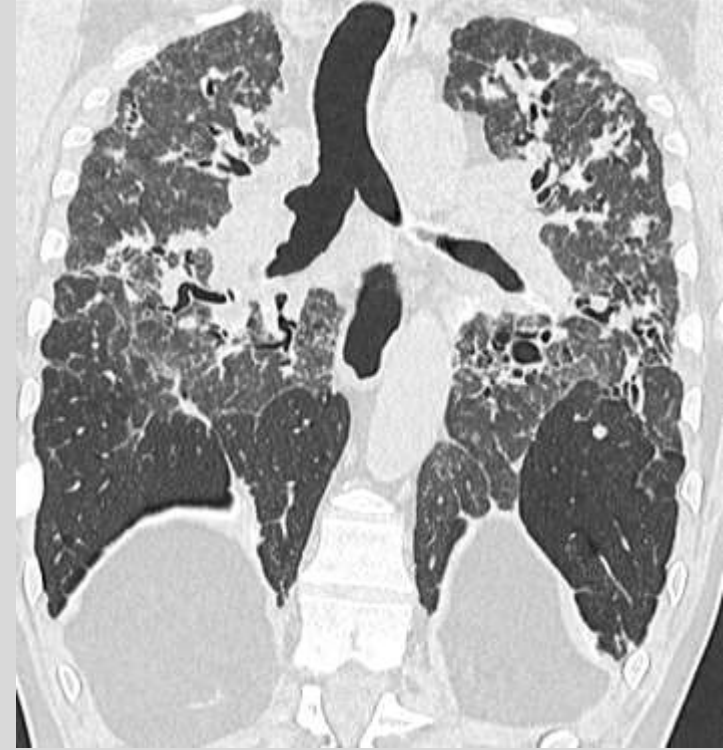
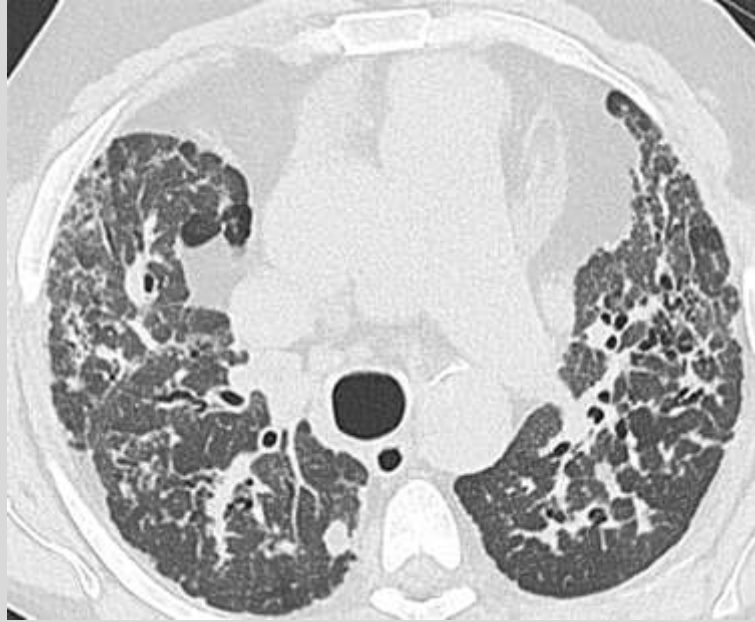


Mou mIP 10 mm



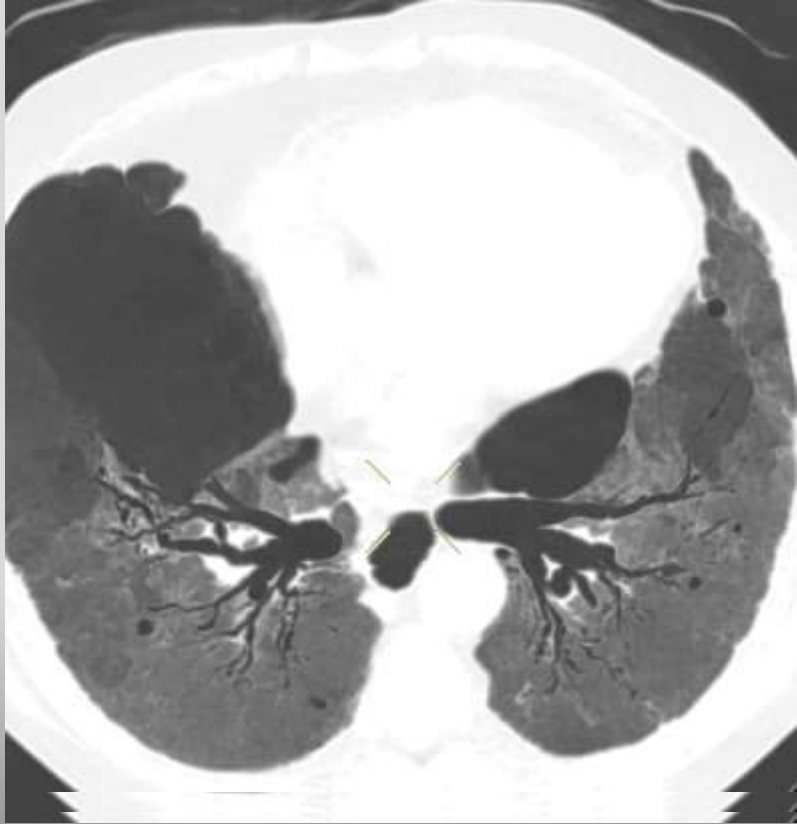
Expiration mou mIP 10 mm

Poumon des boulangers
Bilan pré-greffe pulmonaire



Relatif respect des bases





Développement et validation d'un modèle de diagnostic radiologique pour PHS et score à points model-based

Inclusion:

Cohorte dérivation (Michigan)	n=356	33,9% HP
Cohorte de validation cohort (Lung Tissue Research Consortium)	n=424	15,5% HP
HRCT en inspiration		

Extension de l'atténuation en mosaïque ou air trapping > réticulation
ET

Distribution axiale diffuse

Spécificité élevée

Risque Faux positif HP <10%

VPP diminuée dans un contexte de faible prévalence

Situation 1 : aspect TDM «typique » de PHSf

- Exposition non retrouvée:

⇒ Un LBA lymphocytaire peut suffire si l'exigence diagnostique est modeste

⇒ Dans les autres cas, la BP est à envisager si elle est faisable ?

- Rechercher à nouveau l'exposition avant ++

- Definite : >90% confidence
- high-confidence : 80–89%
- moderate-confidence 70–79%
- low-confidence 51–69% diagnoses.

History of exposure and/or serum IgG testing	Exposure –
No BAL or BAL without lymphocytosis <u>and</u> either no histopathology or indeterminate histopathology	Low confidence
BAL lymphocytosis without histopathology sampling	Moderate confidence
BAL lymphocytosis with indeterminate histopathology	High confidence
Probable HP histopathology	High confidence
Typical HP histopathology	Definite

PHS fibrosante compatible

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*The three-density pattern was formerly called the “headcheese sign.” It is described in detail in Table 4.

**Variable utility of mosaic attenuation
to distinguish fibrotic hypersensitivity
pneumonitis from idiopathic pulmonary
fibrosis**

CT scans inspi et expi 2 lecteurs

n= 102 patients

PIC n=57; PHS chronique n=45

Scoring semiquantitatif pour l'atténuation en mosaïque

Validation sur cohorte externe

PIC n=34; PHS chronique n=28

Augmentation des lobules hypodenses et air trapping selon guidelines (ATS- Fleischner Society)

↑ spécificité pour le diagnostic de PSH chronique avec ↓ sensibilité

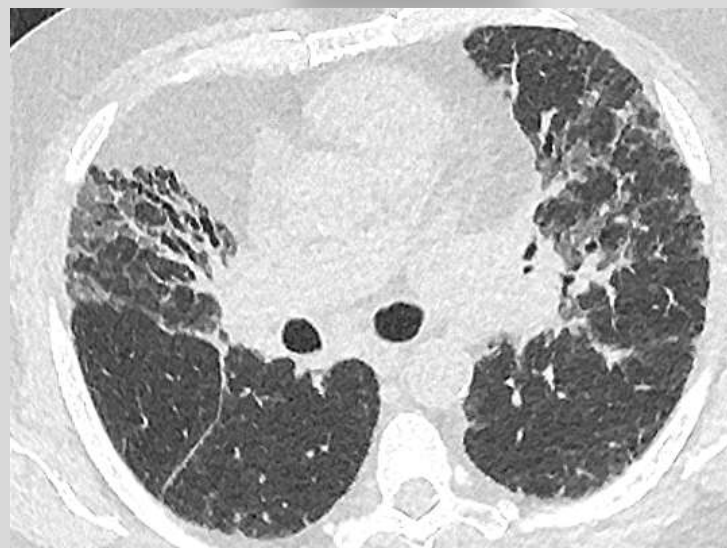
Headcheese sign très spécifique de PHS et inconsistant avec PIC

Atténuation en mosaïque avec lobules hypodenses et air trapping jusque dans 51% des PIC

26/01/17

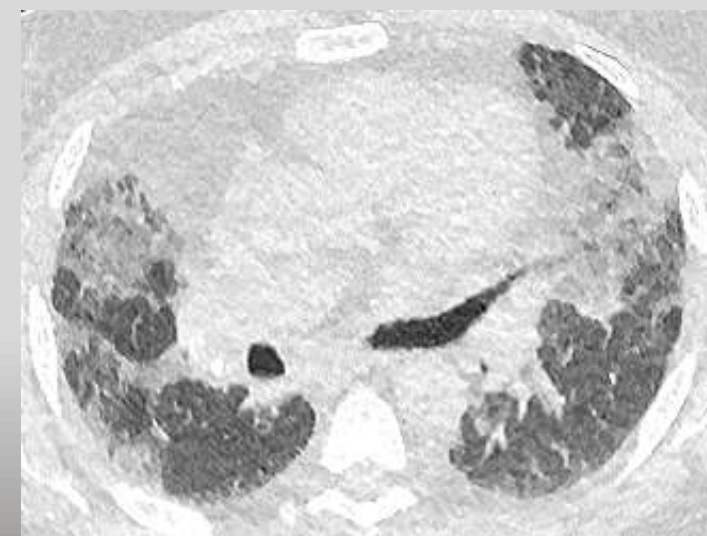
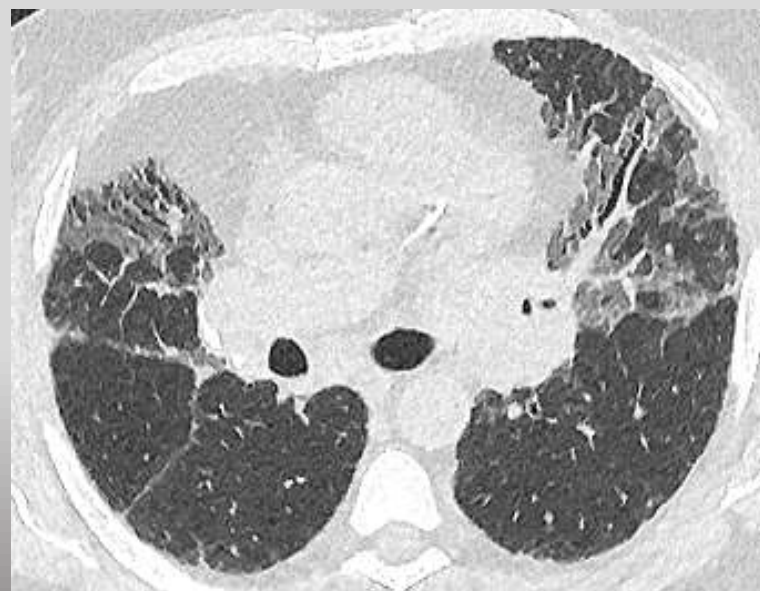
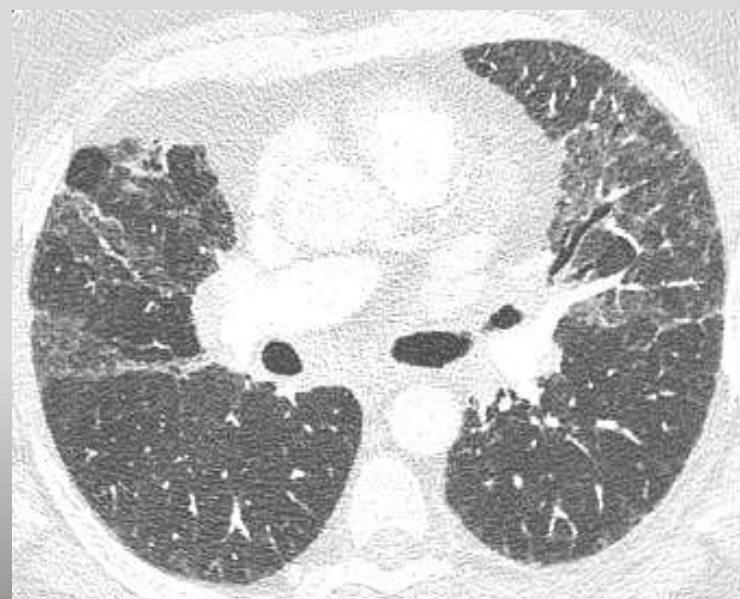


14/06/21

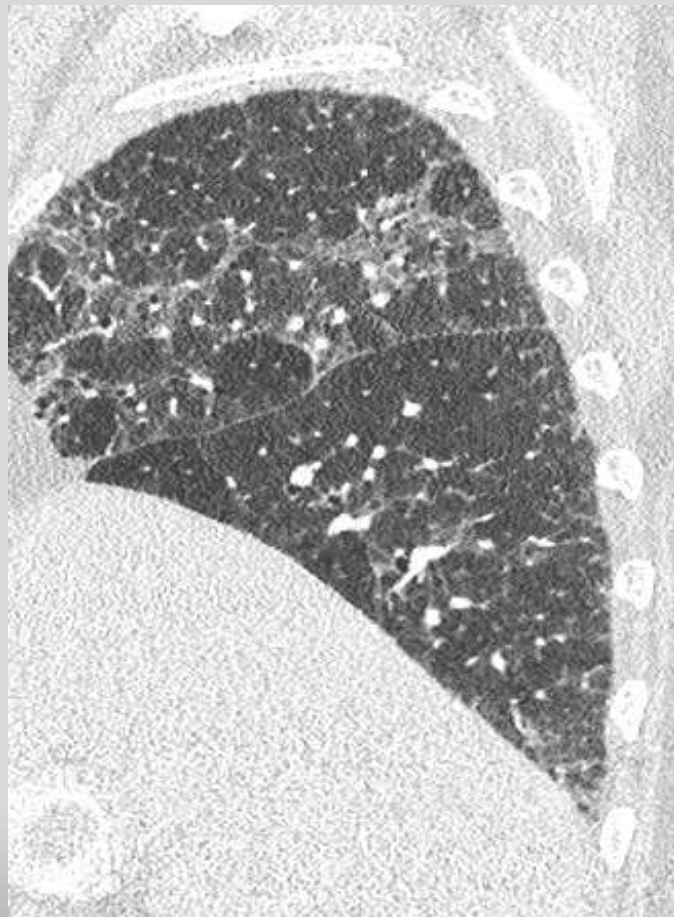


**Prédominance fibreuse en
péribronchovasculaire
Majoration des hyperdensités en
verre dépoli**

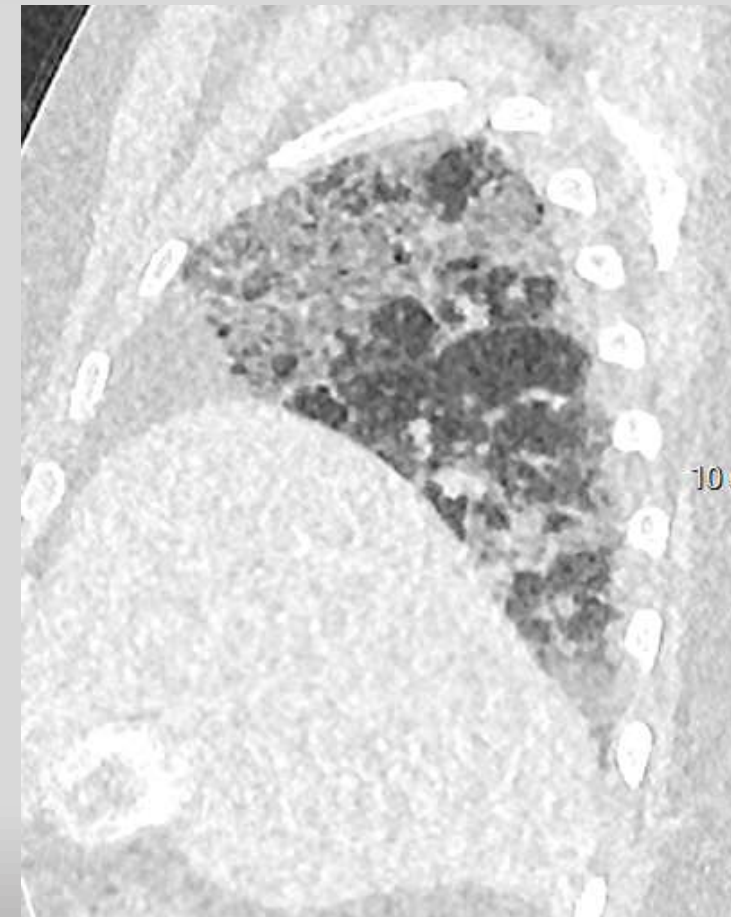
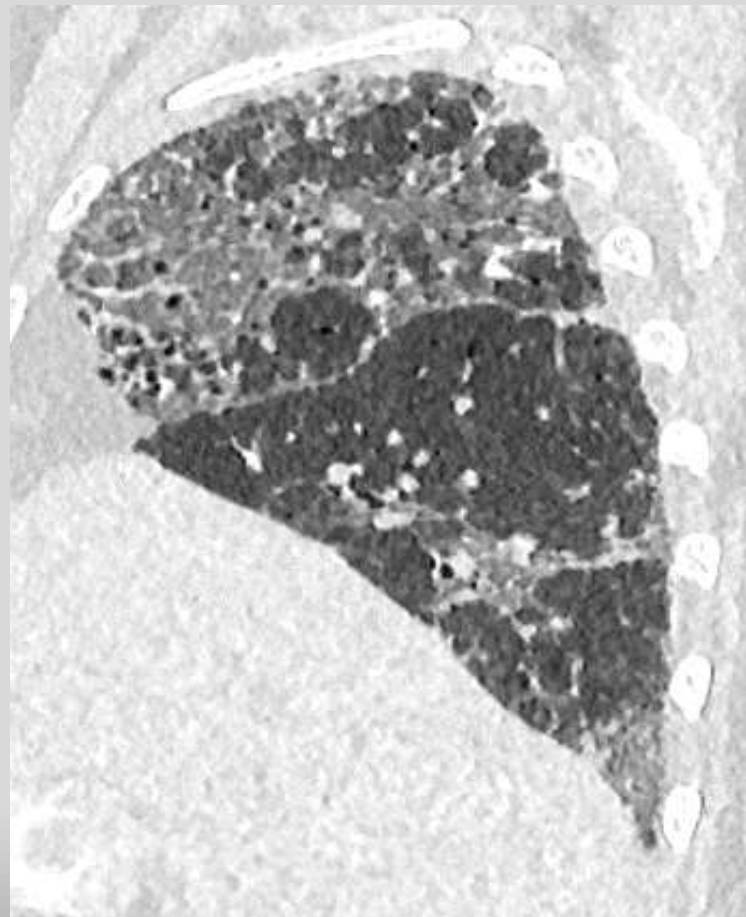
**Classé probable PHS agent causal
indéterminé**



26/01/17



14/06/21



Prédominance fibreuse en péribronchovasculaire et supérieure

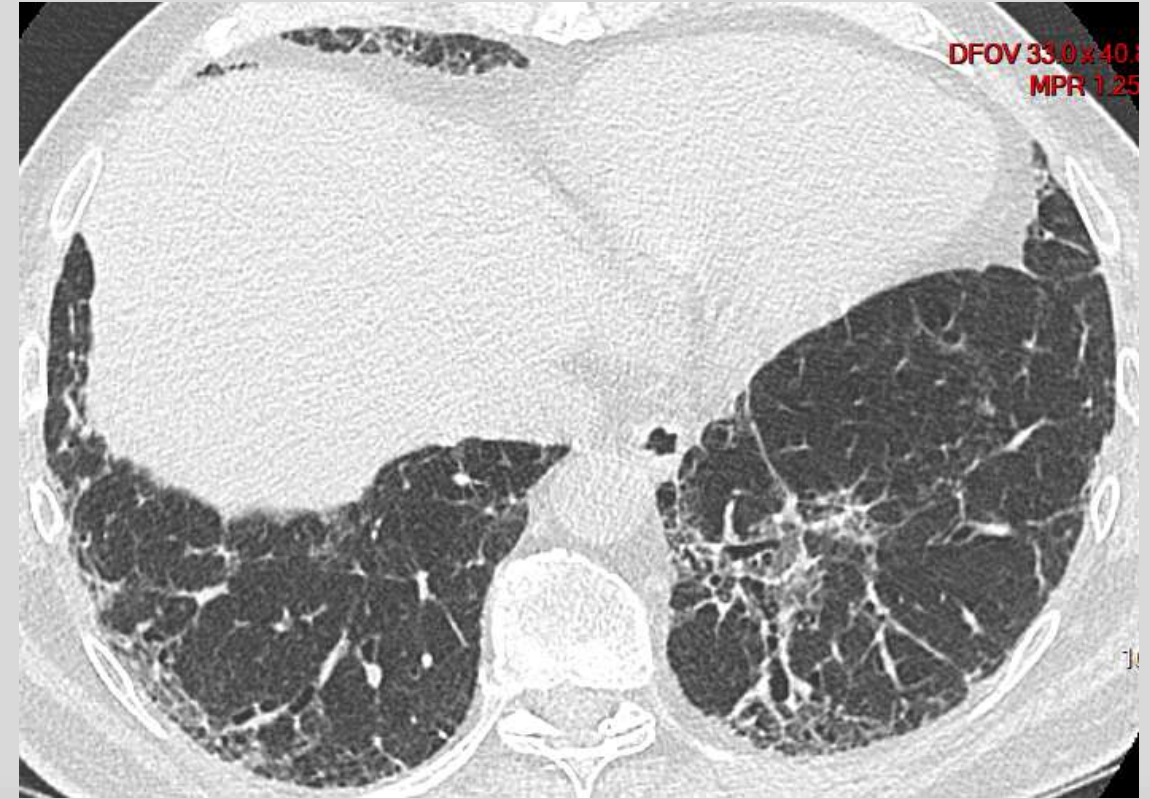
23/04/15



Prédominance fibreuse en péribronchovasculaire et sous-pleurale

Poumon fermier

03/02/20



Noter la perte volume

PHS fibrosante indéterminée

Table 6. Chest HRCT Scan Features of the Fibrotic HP Pattern

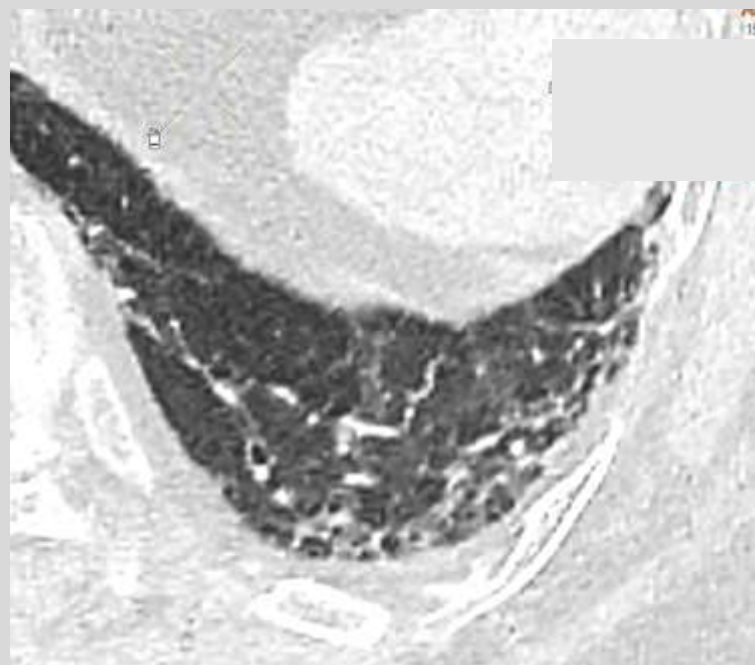
HRCT Pattern	Typical HP	Compatible with HP	Indeterminate for HP
Description	The “typical HP” pattern is suggestive of a diagnosis of HP. It requires a) an HRCT pattern of lung fibrosis (as listed below) in one of the distributions and b) at least one abnormality that is indicative of small airway disease	“Compatible-with-HP” patterns exist when the HRCT pattern and/or distribution of lung fibrosis varies from that of the typical HP pattern; the variant fibrosis should be accompanied by signs of small airway disease	The “indeterminate-for-HP” pattern exists when the HRCT is neither suggestive nor compatible with a typical and probable HP pattern
Relevant radiological findings	HRCT abnormalities indicative of lung fibrosis are most commonly composed of irregular linear opacities/coarse reticulation with lung distortion; traction bronchiectasis and honeycombing may be present but do not predominate The distribution of fibrosis may be: • Random both axially and craniocaudally or • Mid lung zone–predominant or • Relatively spared in the lower lung zones HRCT abnormalities indicative of small airway disease: • Ill-defined, centrilobular nodules and/or GGOs • Mosaic attenuation, three-density pattern,* and/or air trapping (<i>often in a lobular distribution</i>)	Variant patterns of lung fibrosis: • UIP pattern: basal and subpleural distribution of honeycombing with/without traction bronchiectasis (<i>per 2018 diagnosis of IPF guidelines</i> [20]) • Extensive GGOs with superimposed subtle features of lung fibrosis Variant (predominant) distributions of lung fibrosis: • Axial: peribronchovascular, subpleural areas • Craniocaudal: upper lung zones HRCT abnormalities indicative of small airway disease: • Ill-defined centrilobular nodules, or • Three-density pattern* and/or air trapping	One patterns (i.e., not accompanied by other findings suggestive of HP) of: • UIP pattern (<i>as per 2018 IPF diagnosis guidelines</i> [20]) • Probable UIP pattern (<i>as per 2018 IPF diagnosis guidelines</i> [20]) • Indeterminate pattern for UIP (<i>as per 2018 IPF diagnosis guidelines</i> [20]) • Fibrotic NSIP pattern • Organizing pneumonia–like pattern • Truly indeterminate HRCT pattern

Definition of abbreviations: GGO = ground-glass opacity; HP = hypersensitivity pneumonitis; HRCT = high-resolution computed tomography;

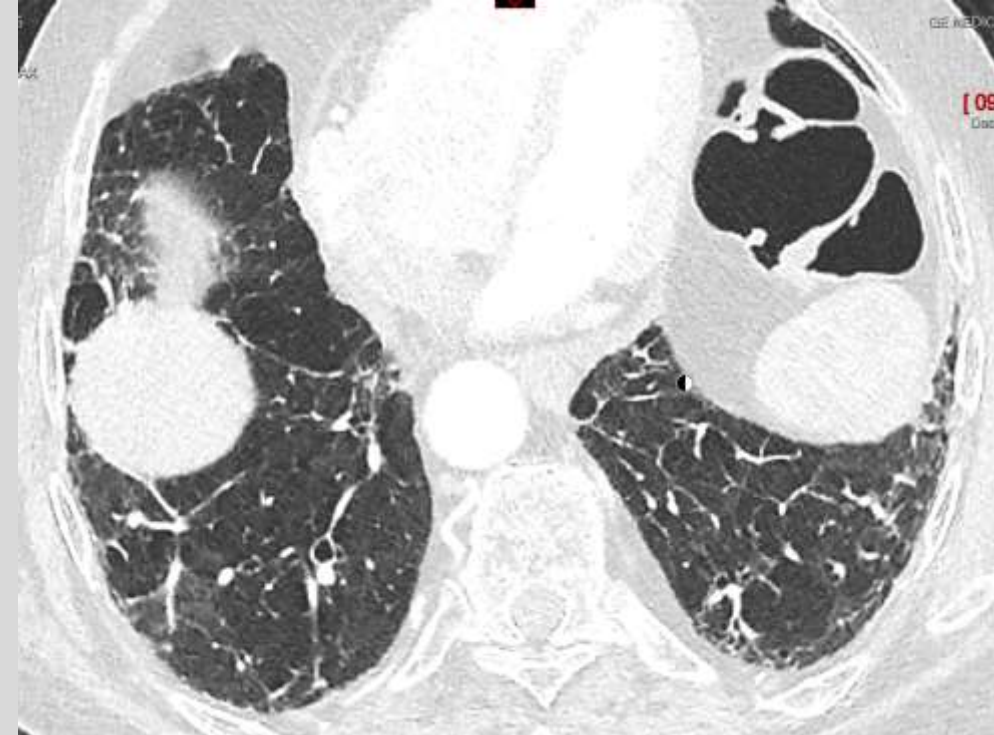
IPF = idiopathic pulmonary fibrosis; NSIP = nonspecific interstitial pneumonia; UIP = usual interstitial pneumonia.

Rarely, fibrotic HP may be seen 1) as a component of combined pulmonary fibrosis and emphysema or pleuroparenchymal fibroelastosis with emphysema, 2) as a pure emphysematous form of HP, or 3) in acute exacerbation.

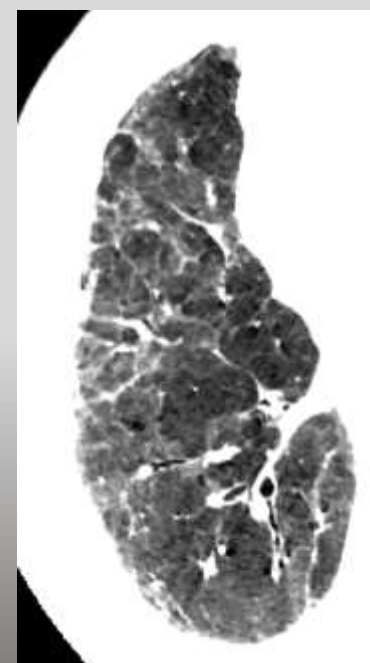
*The three-density pattern was formerly called the “headcheese sign.” It is described in detail in Table 4.



Probable UIP
PHS ?



Hernie hiatale



16/12/19



13/09/21



PHS le plus probable
DD NSIP
Prednisone Certain bénéfice

En co-association
avec lobules
hyperclairs

Situation 1 : aspect TDM «typique » de PHSf

- Exposition retrouvée:

- LBA lymphocytaire
⇒ certitude diagnostique par BP est certainement futile

- LBA non lymphocytaire
⇒ BP peut s'avérer nécessaire

- Definite : >90% confidence
- high-confidence : 80–89%
- moderate-confidence 70–79%
- low-confidence 51–69% diagnoses.

History of exposure and/or serum IgG testing	Exposure +
No BAL or BAL without lymphocytosis <u>and</u> either no histopathology or indeterminate histopathology	Moderate confidence
BAL lymphocytosis without histopathology sampling	High confidence
BAL lymphocytosis with indeterminate histopathology	Definite
Probable HP histopathology	Definite
Typical HP histopathology	Definite

Situation 2 : aspect TDM «compatible» ou « indéterminé » de PHSf

	Compatible with HP		Indeterminate for HP	
History of exposure and/or serum IgG testing	Exposure +	Exposure –	Exposure +	Exposure –
No BAL or BAL without lymphocytosis <u>and</u> either no histopathology or indeterminate histopathology	Low confidence	Not excluded	Not excluded	Not Excluded
BAL lymphocytosis without histopathology sampling	Moderate confidence	Low confidence	Low confidence	Not excluded

- Definite : >90% confidence
- high-confidence : 80–89%
- moderate-confidence 70–79%
- low-confidence 51–69% diagnoses.

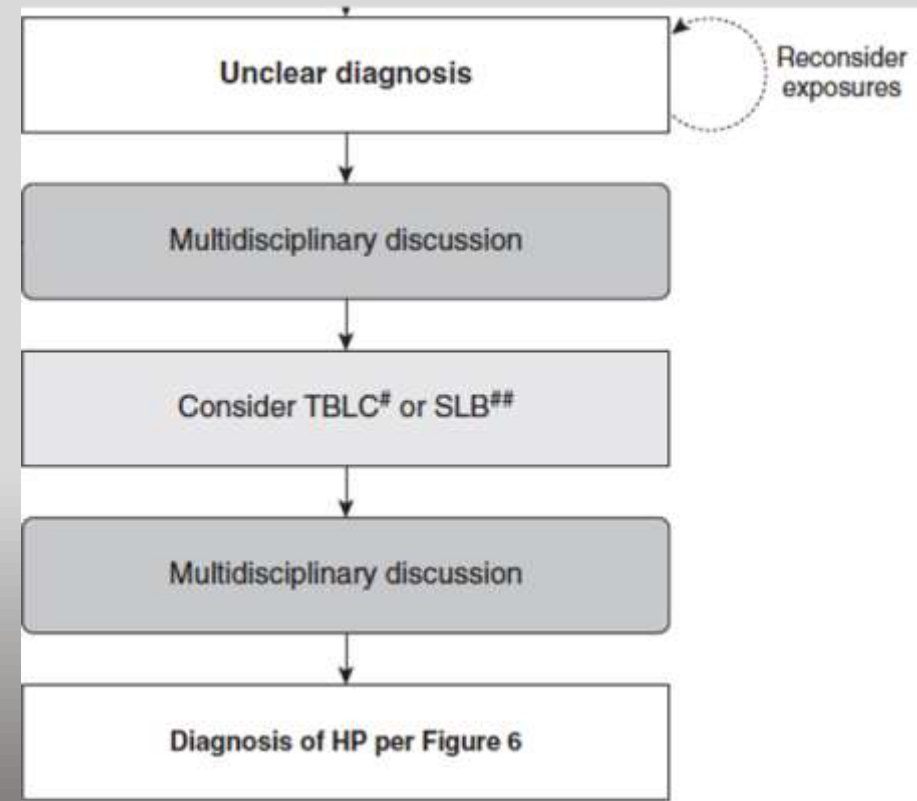
Situation 2 : aspect TDM «compatible» ou « indéterminé » de PHSf

- La BP sera souvent nécessaire

	Compatible with HP		Indeterminate for HP	
History of exposure and/or serum IgG testing	Exposure +	Exposure –	Exposure +	Exposure –
No BAL or BAL without lymphocytosis and either no histopathology or indeterminate histopathology	Low confidence	Not excluded	Not excluded	Not Excluded
BAL lymphocytosis without histopathology sampling	Moderate confidence	Low confidence	Low confidence	Not excluded

Tableau 5. Principales contre-indications à la biopsie pulmonaire vidéo-chirurgicale

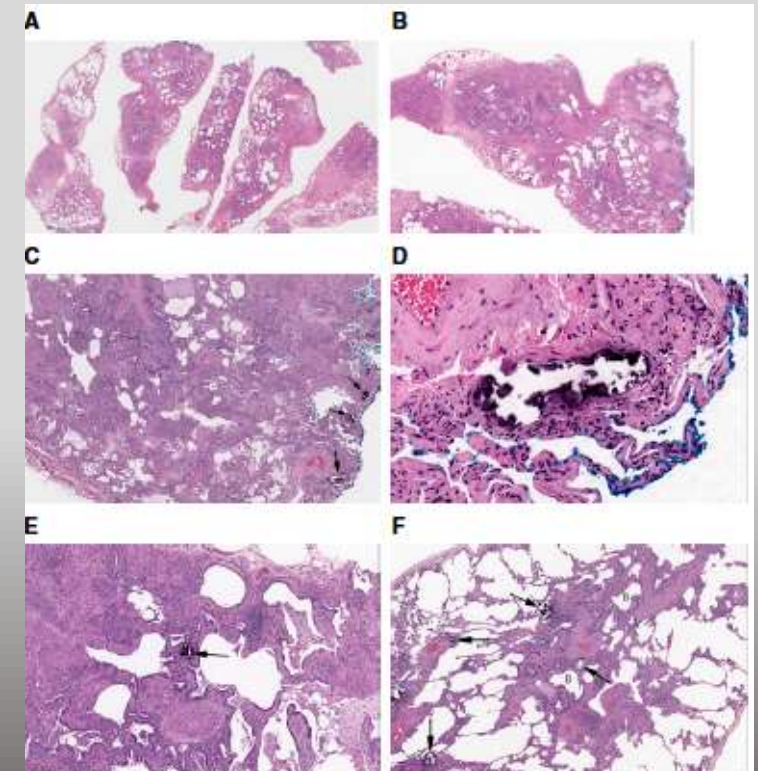
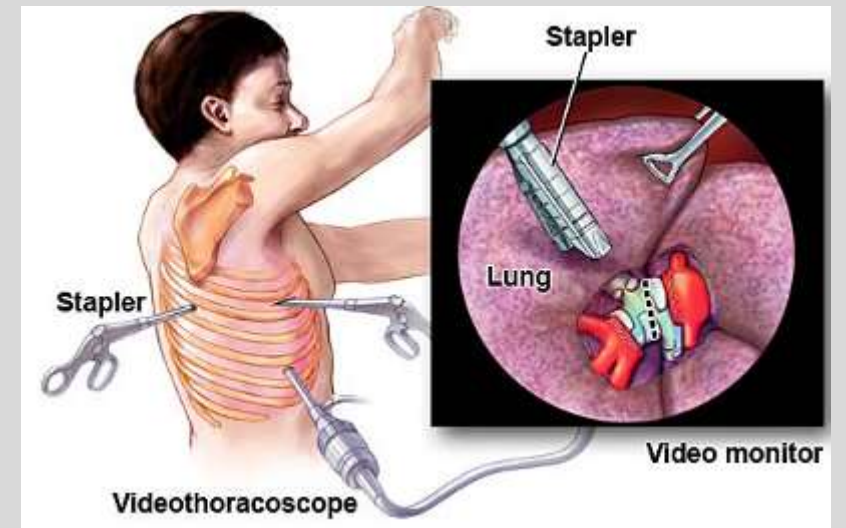
- Âge physiologique > 75 ans
- Aggravation rapide de la maladie
- Faible réserve respiratoire (CVF < 60-70 %, DLco < 35-40 %)
- Oxygénothérapie de longue durée
- Hypertension pulmonaire
- Comorbidités importantes ou multiples
- Immunodépression



Biopsie Pulmonaire

Table 7. Histopathological Criteria for the Diagnosis of HP (Other than "Hot-Tub Lung")

HP	Probable HP	Indeterminate for HP
Fibrotic HP[‡] Typical histopathological features of fibrotic HP; 1 or 2 and 3 in at least one biopsy site: - Chronic fibrosing interstitial pneumonia - Architectural distortion, fibroblast foci ± subpleural honeycombing - Fibrotic NSIP-like[§] pattern - Airway-centered fibrosis ± Peribronchiolar metaplasia ± Bridging fibrosis	Both of the following features (1 or 2 from first column) in at least one biopsy site: - Chronic fibrosing interstitial pneumonia - Architectural distortion, fibroblast foci ± subpleural honeycombing - Fibrotic NSIP-like pattern - Airway-centered fibrosis ± Peribronchiolar metaplasia ± Bridging fibrosis ± Cellular interstitial pneumonia ± Organizing pneumonia pattern ± Cellular bronchiolitis and Absence of features in any biopsy site to suggest an alternative diagnosis	Either one of the following features in at least one biopsy site: - Chronic fibrosing interstitial pneumonia - Architectural distortion, fibroblast foci ± honeycombing - Fibrotic NSIP-like pattern ± Cellular interstitial pneumonia ± Cellular bronchiolitis ± Organizing pneumonia pattern and Absence of features in any biopsy site to suggest an alternative diagnosis - Plasma cells > lymphs - Extensive lymphoid hyperplasia - Extensive well-formed sarcoidal granulomas and/or necrotizing granulomas - Aspirated particulates
- Poorly formed nonnecrotizing granulomas[†] ± Cellular interstitial pneumonia ± Cellular bronchiolitis ± Organizing pneumonia pattern and Absence of features in any biopsy site to suggest an alternative diagnosis - Plasma cells > lymphs - Extensive lymphoid hyperplasia - Extensive well-formed sarcoidal granulomas and/or necrotizing granulomas - Aspirated particulates		

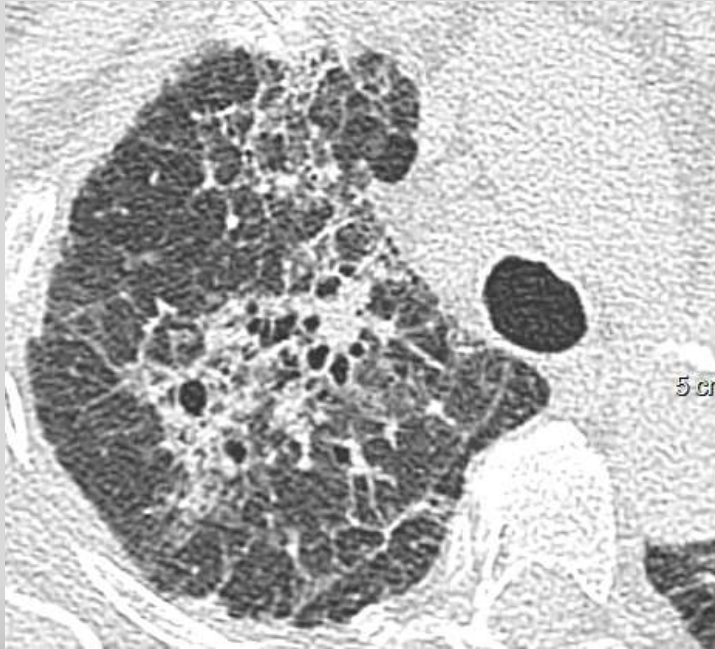


Situation 2 : aspect TDM «compatible» ou « indéterminé » de PHSf : BP effectuée

	Compatible with HP		Indeterminate for HP	
History of exposure and/or serum IgG testing	Exposure +	Exposure –	Exposure +	Exposure –
No BAL or BAL without lymphocytosis <u>and</u> either no histopathology or indeterminate histopathology	Low confidence	Not excluded	Not excluded	Not Excluded
BAL lymphocytosis without histopathology sampling	Moderate confidence	Low confidence	Low confidence	Not excluded
BAL lymphocytosis with indeterminate histopathology	Moderate confidence	Moderate confidence	Low confidence	Not excluded
Probable HP histopathology	High confidence	Moderate confidence	Moderate confidence	Low confidence
Typical HP histopathology	Definite	Definite	Definite	High confidence*

Limitations- Questions-Remarques

**Quelle est la caractéristique dominante?
Pattern? Distribution axiale? CC?**

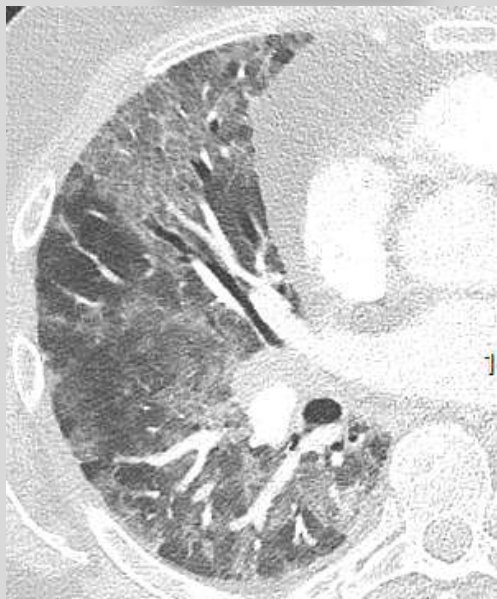


Réticulations irrégulières



Périfonchovasculaire et sous-pleurale

24/11/19



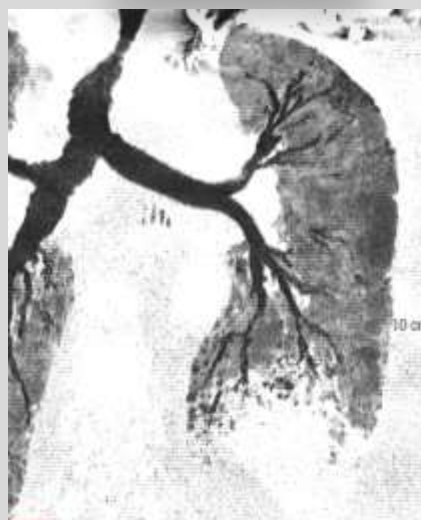
20/03/20



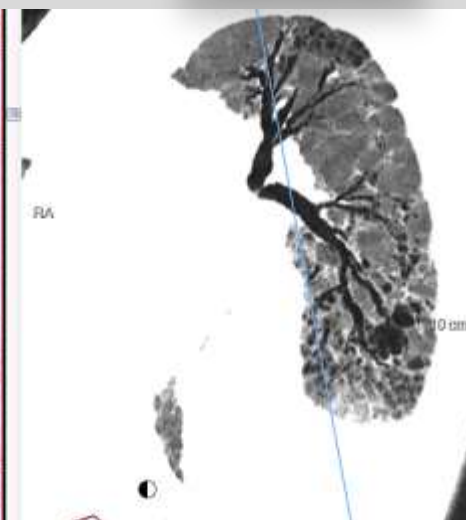
Pneumonie à CMV

Attention aux diagnostics différentiels !

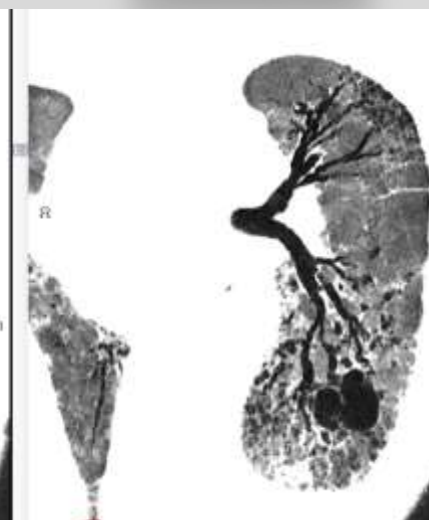
24/11/19



20/03/20

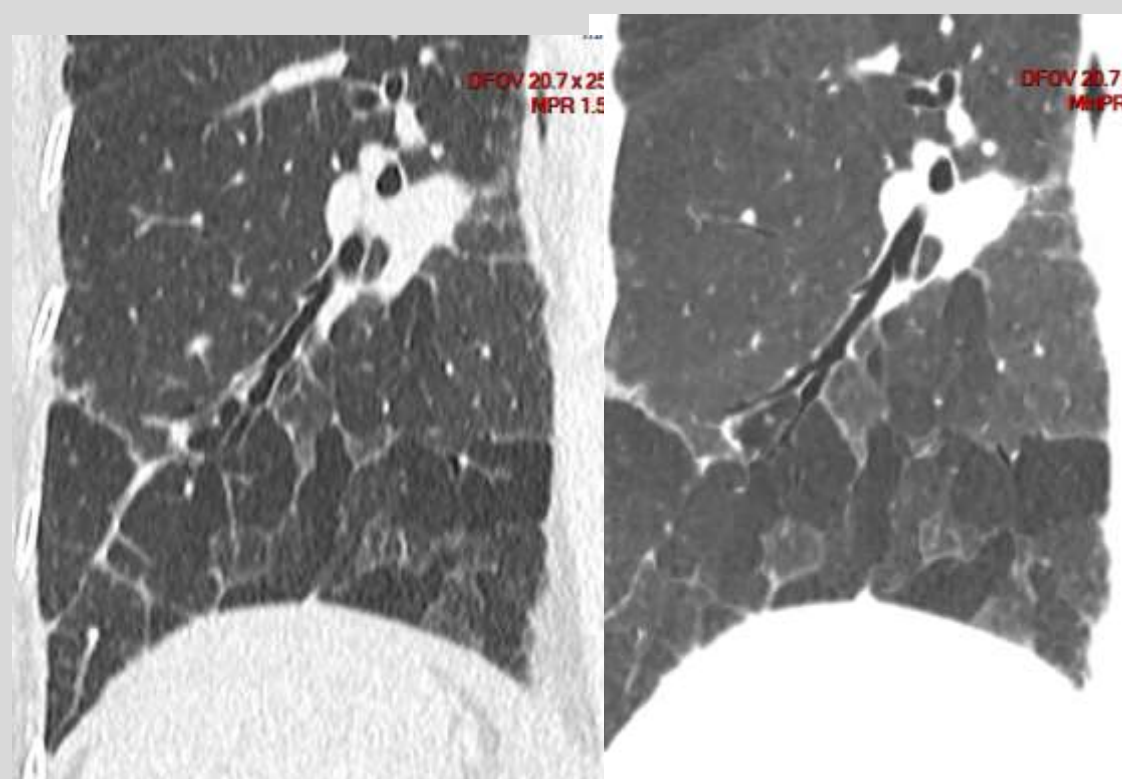


14/09/20



PHS chronique rapidement progressive depuis 2014

O2 au long cours.
Prednisone / Azathioprine / Nintedanib



Exposition environnementale
avec précipitines positives pour *A. fumigatus*, *M. faeni*, *P. casei* et humidificateur

Eviction complète des antigènes incriminés
Arrêt du traitement IS par la patiente
Progression fibrose avec perte de volume du LIG

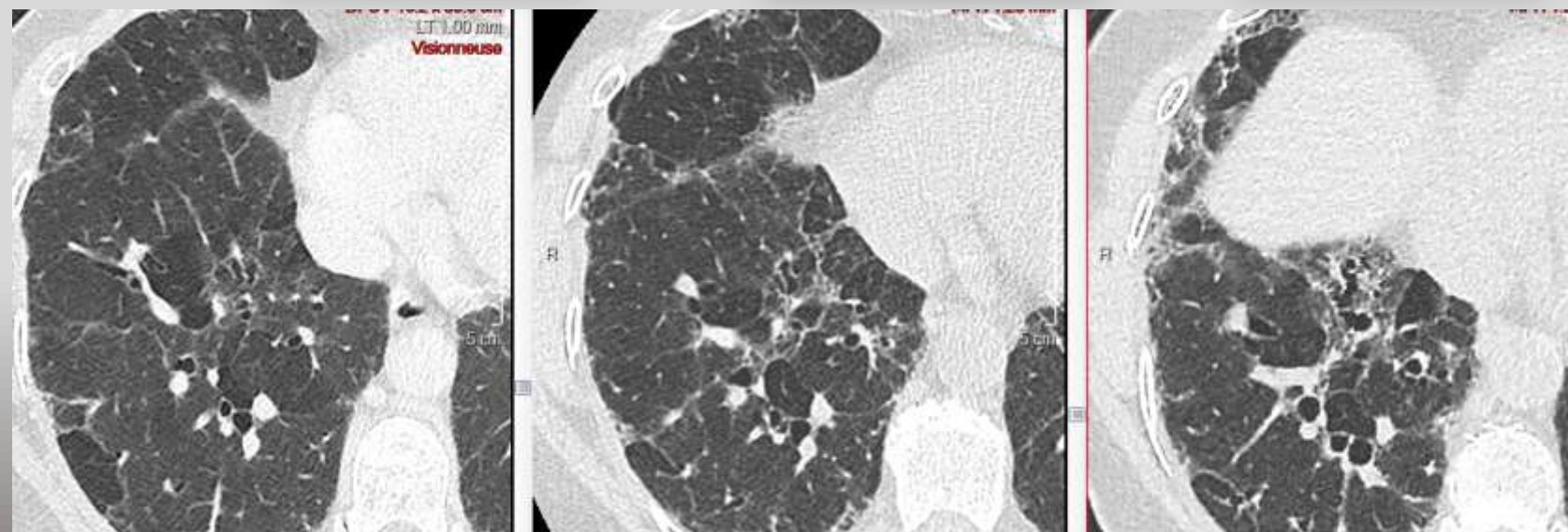
12/11/13

17/12/18

07/05/21

12/11/13

**A quel moment
est effectuée
imagerie ?**



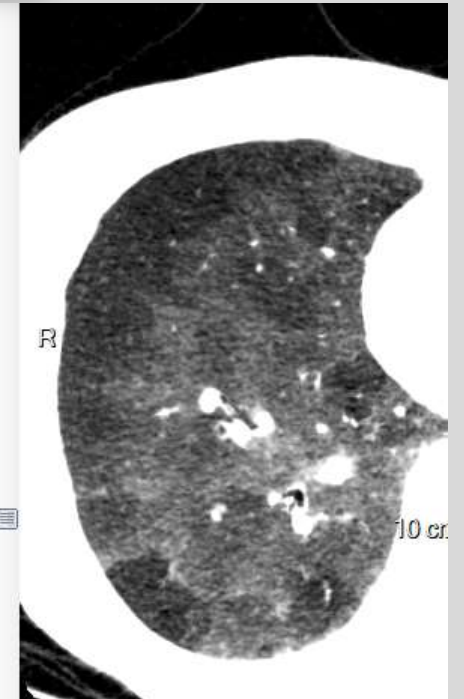
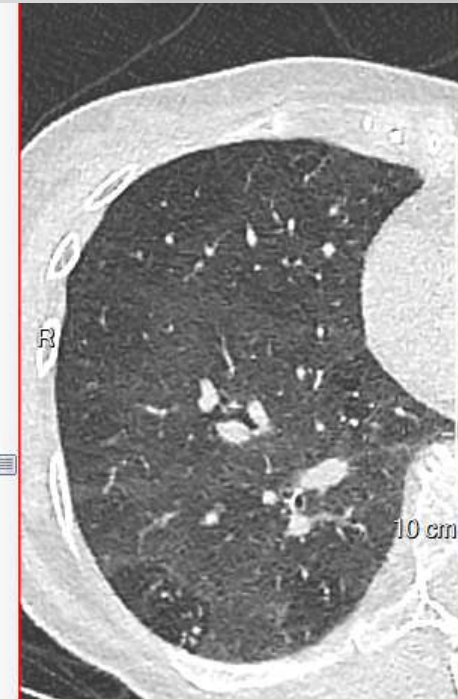
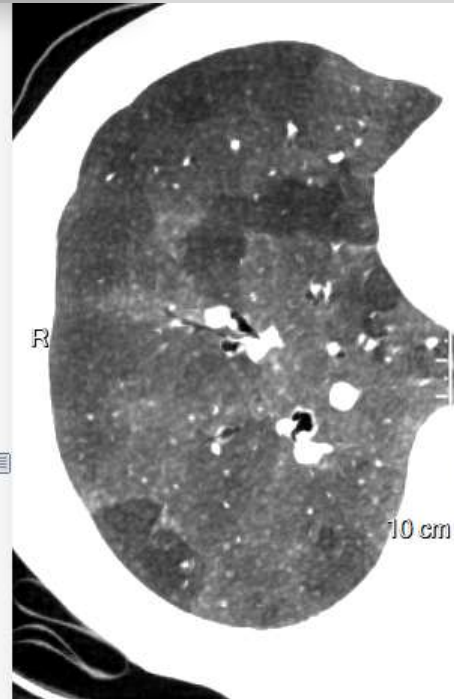
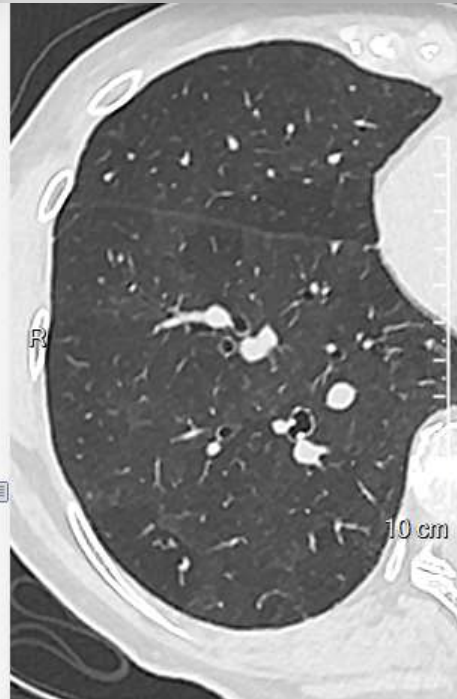
PHS Aiguë humidificateurs et jacuzzi
Précipitines pour *M. faeni* et humidificateur positives

Phase aiguë

Asymptomatique

Expiration
?

Signification ?



Lung

Médiastin

M et mIP

L expi

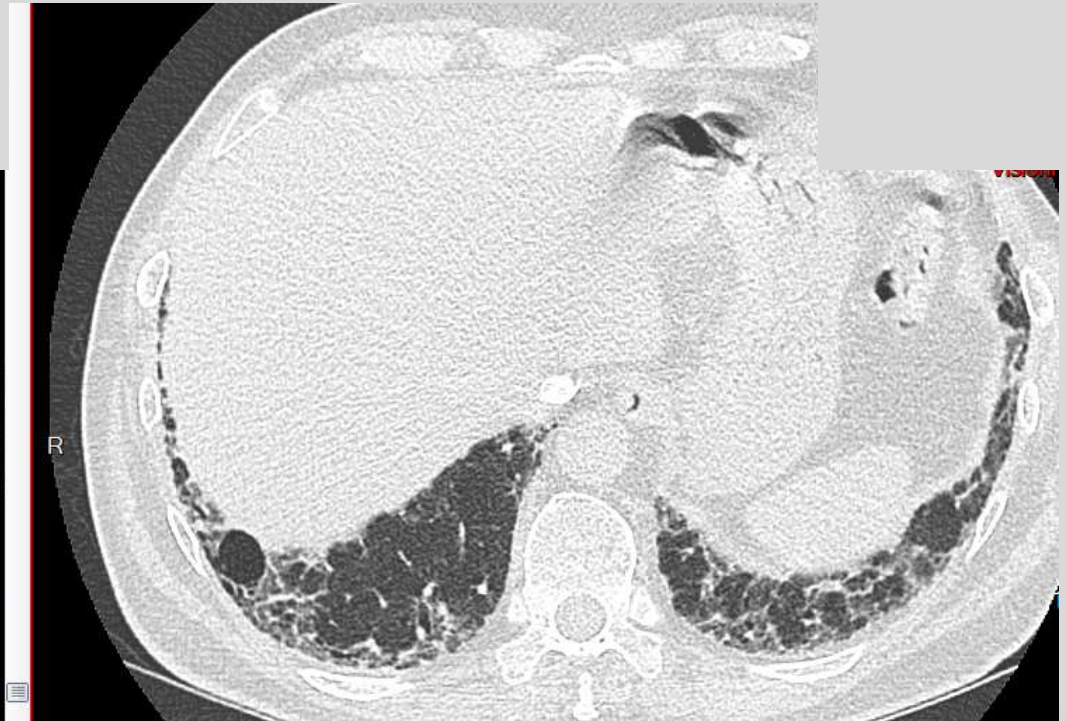
M expi mIP

Kystes

20/08/18



03/02/20



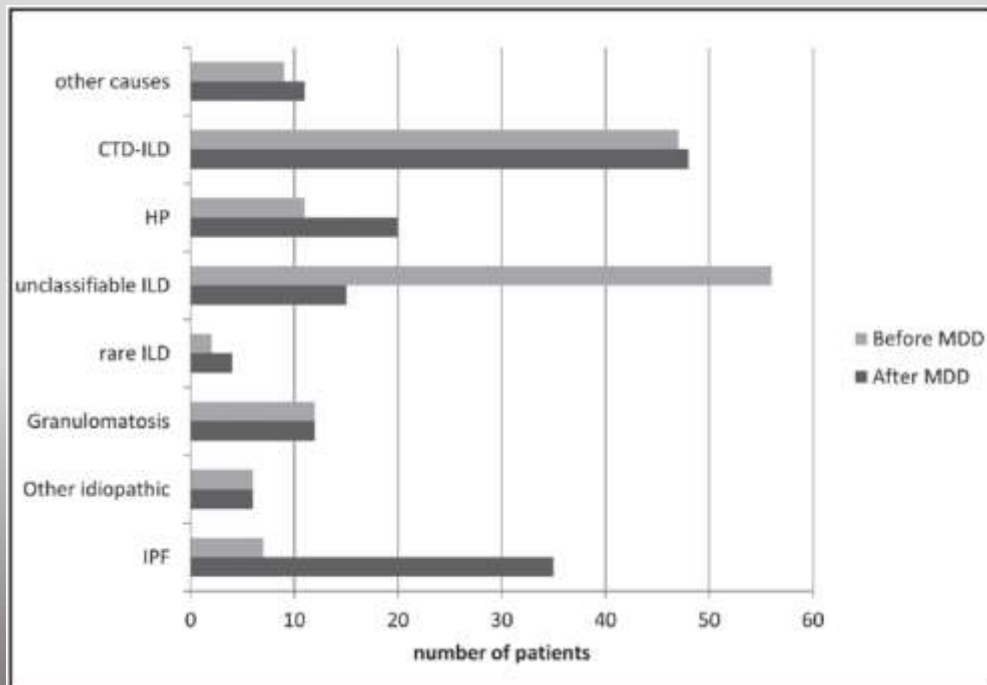
PHS à *Saccharopolyspora rectivirgula*
"poumon de fermier" en 1999
fin d'exposition en 2001

Situation 2 : aspect TDM «compatible» ou « indéterminé » de PHSf : BP non effectuée

MULTIDISCIPLINARY MANAGEMENT OF INTERSTITIAL LUNG DISEASES: A REAL-LIFE STUDY

Caroline Biglia¹, Benoit Gbaze², Gregory Reyblier^{1,2}, Sandra Koenig¹, Halil Yildiz³, Valérie Lacroix⁴, Farah Tamirou⁵, Delphine Hoton⁶, Thierry Pieters¹, Antoine Froidure^{1,2}

Sarcoidosis Vasc Diffuse Lung Dis 2019; 36 (2): 108-115



Unclassifiable interstitial lung disease: from phenotyping to possible treatments

Sabina A. Guler^{a,b,c} and Christopher J. Ryerson^{a,b}

Curr Opin Pulm Med 2018, 24:461-468

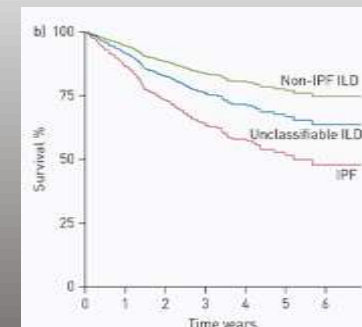
Table 1. Reasons for unclassifiable interstitial lung disease

	Clinical [11,12 ^a]	Radiological [19 ^a]	Pathological [20 ^a]	Multidisciplinary discussion [11,12 ^a ,21]
Incomplete evaluation	Unable to obtain adequate history (e.g., exposures)	Not available HRCT quality insufficient	No biopsy performed (e.g., unfavorable risk-benefit ratio, patient preference) Insufficient biopsy quality (too small, damaged, nonoptimal sampling location)	Not available Difficult interpretation of poor quality diagnostic material
Overlapping findings	Multiple risk factors predisposing for different specific ILDs	Overlap with non-ILD features (e.g., cardiac failure, infection) Acute exacerbation Features of different specific ILDs	Overlapping histological features	Discrepant clinical, radiological, and pathological features Discrepant interpretation of information by members
Nonspecific findings	Stable disease, mild symptoms	Indeterminate for UIP Prior treatment (e.g., corticosteroids)	Only advanced interstitial fibrosis Prior treatment (e.g., corticosteroids)	Poorly classifiable findings

Prevalence and prognosis of unclassifiable interstitial lung disease

Christopher J. Ryerson¹, Thomas H. Urbania¹, Luca Richeldi², Joshua J. Mooney³, Joyce S. Lee⁴, Kirk D. Jones⁵, Brett M. Elicker⁶, Laura L. Kott⁷, Talmadge E. King Jr¹, Paul J. Walters⁸ and Harold R. Colvard¹

The Journal of Thoracic Medicine and Lung Transplantation



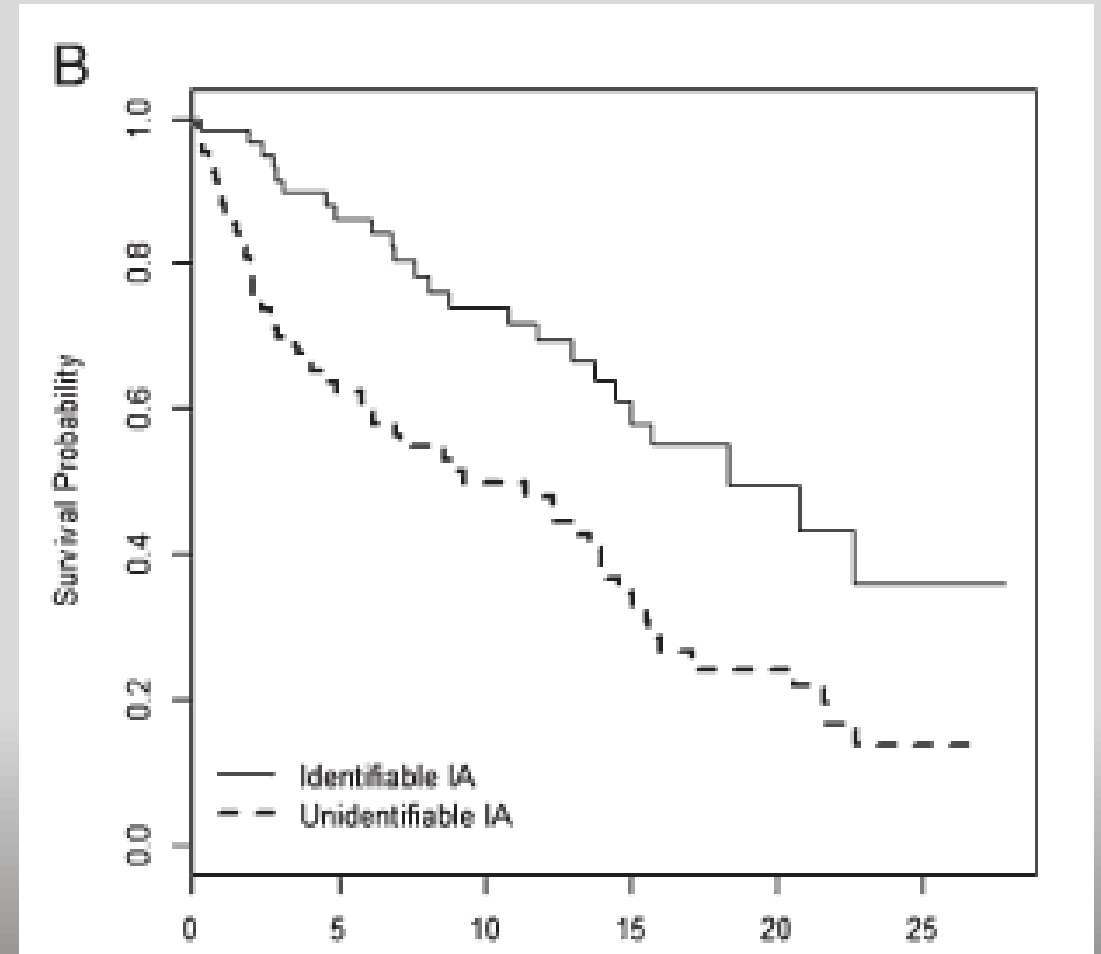
Prise en charge

1- Trouver l'antigène et l'évincer

Identifying an Inciting Antigen Is Associated With Improved Survival in Patients With Chronic Hypersensitivity Pneumonitis

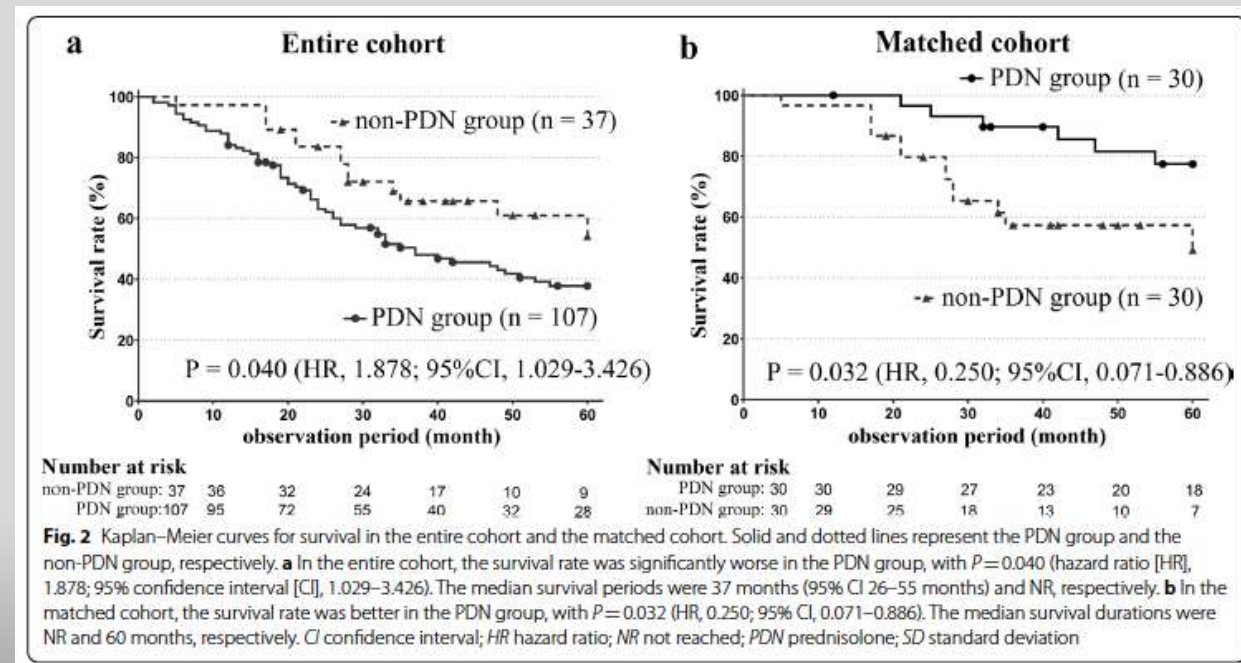
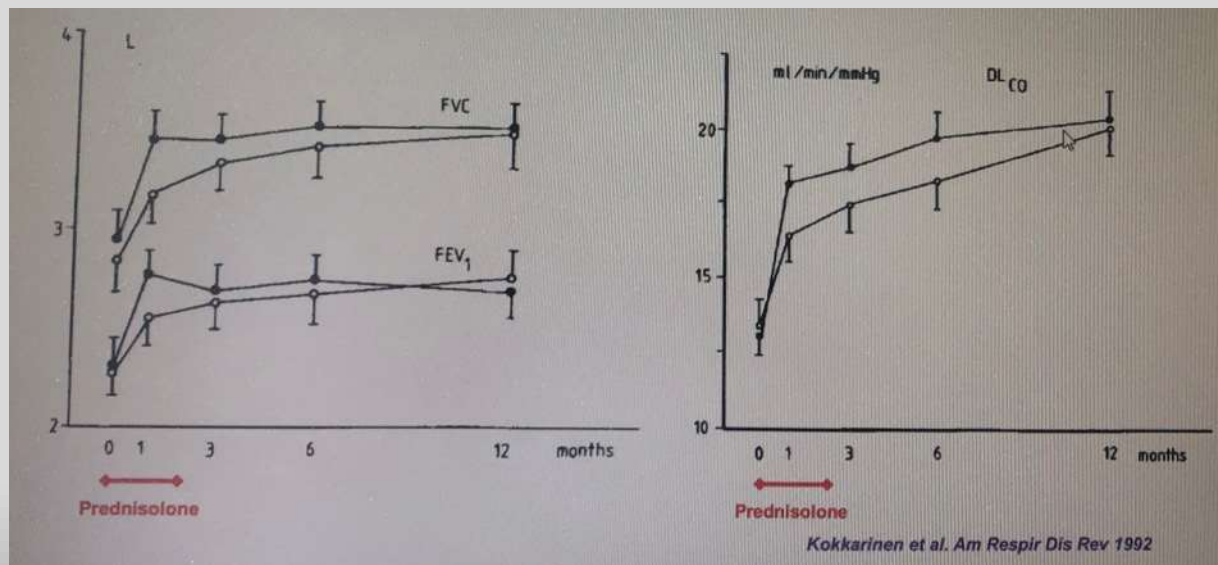
Evans R. Fernández Pérez, MD, FCCP; Jeffrey J. Swigris, DO, FCCP; Anna V. Forssén, MS; Olga Tourin, MD; Joshua J. Solomon, MD, FCCP; Tristan J. Huie, MD, FCCP; Amy L. Olson, MD, MSPH; and Kevin K. Brown, MD, FCCP

CHEST 2013; 144(5):1644–1651



Prise en charge

2- Traiter par corticothérapie



Prise en charge

3- Ajouter des IS ?

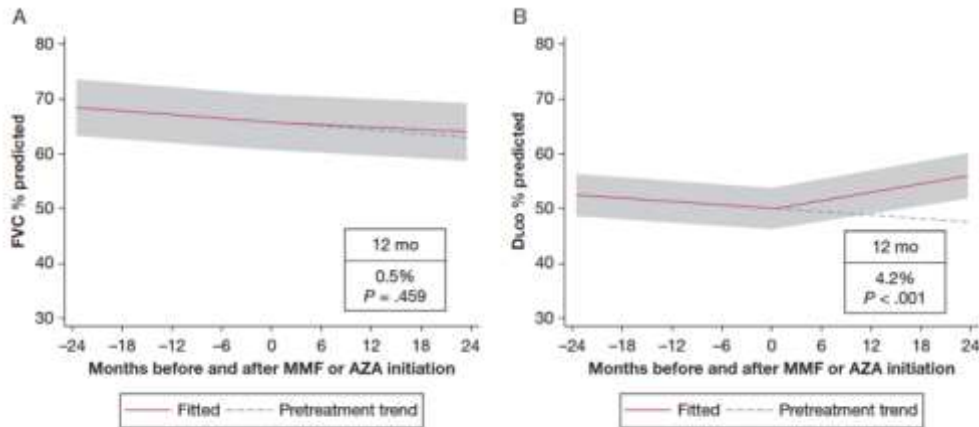


Figure 2 – Mixed-effects model estimates for FVC % predicted and DLCO % predicted before and after initiation of mycophenolate or azathioprine. The gray shading indicates the 95% CI. DLCO = diffusion capacity of the lung for carbon monoxide. See Figure 1 legend for expansion of other abbreviations.

CHEST 2017; 151(3):619-625

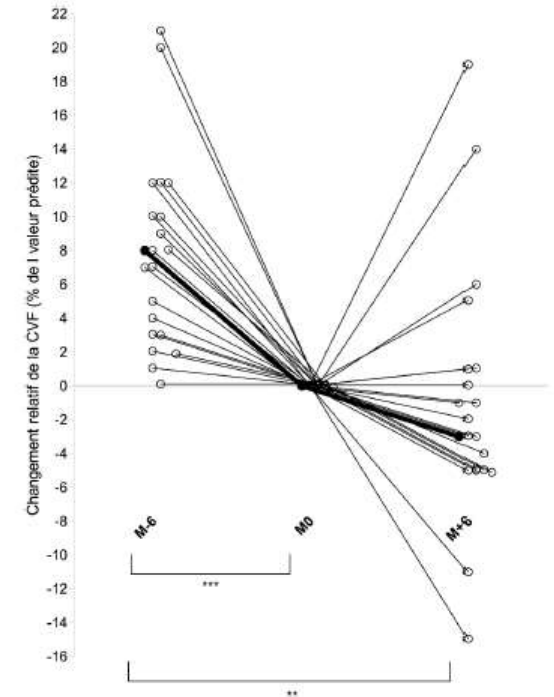
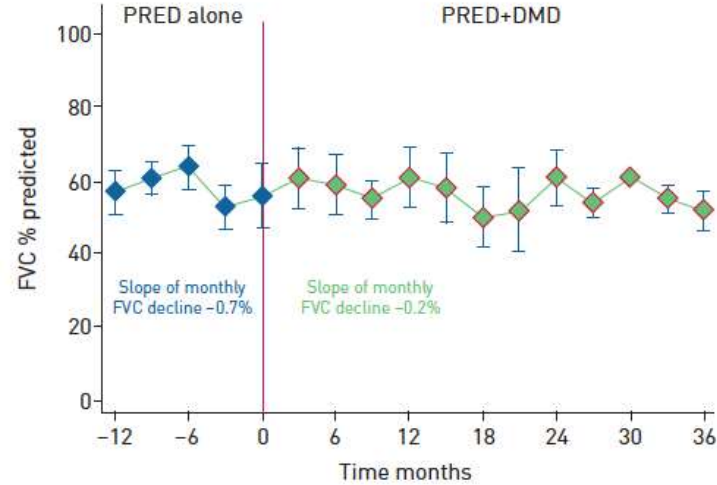
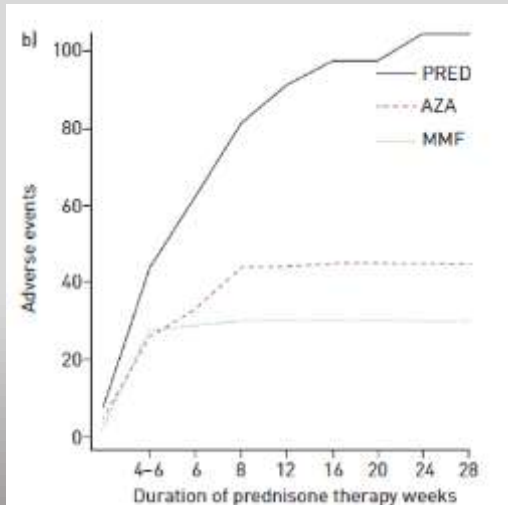


Fig. 1. Relative change in FVC (% of predicted value), 6 months before and after the introduction of rituximab (n = 20). The median value is represented by the bold line. ** and ***: p < 0.01 and < 0.001, respectively.



ERJ Open Res 2017; 3: 00016-2017

Respiratory Medicine 172 (2020) 106146

Traitement des Fibroses Pulmonaires hors FPI

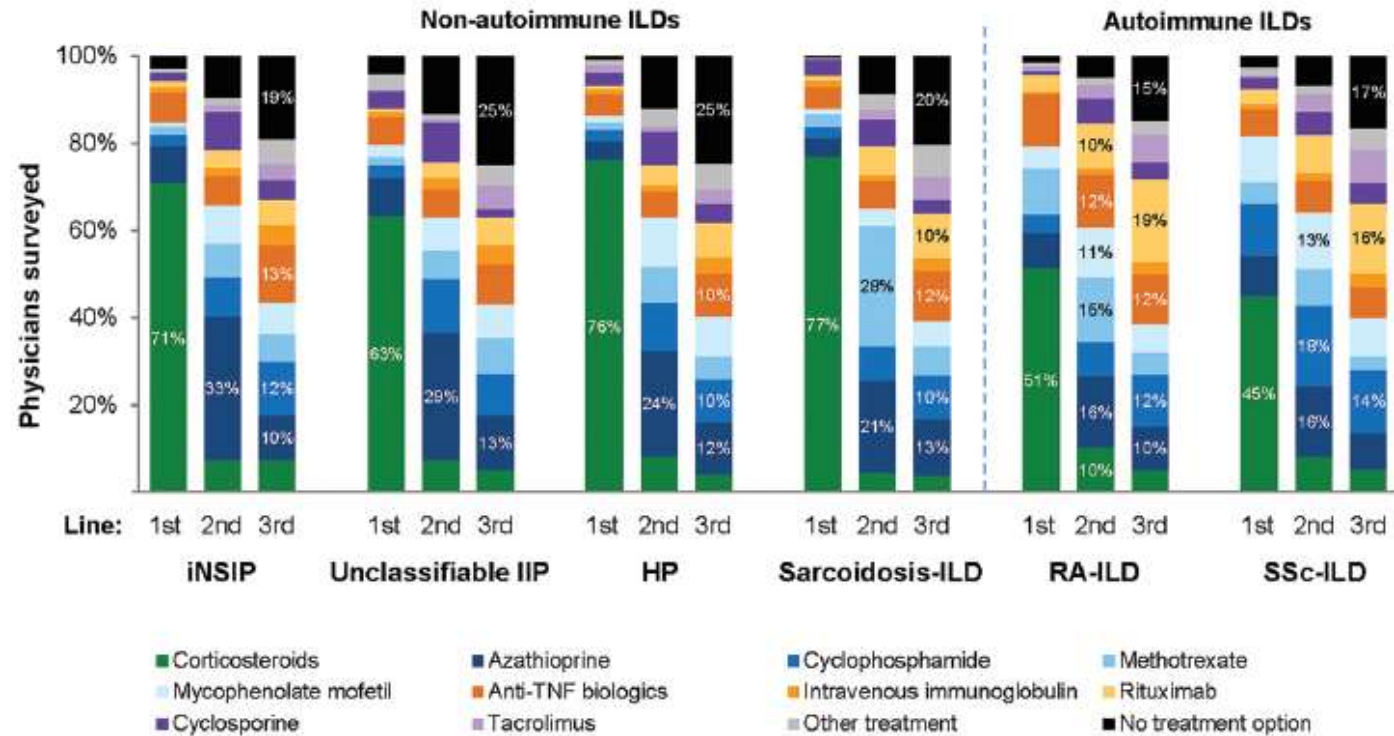


Figure 7. Agents used as first-, second- and third-line treatments for fibrotic ILDs. Data from online survey of physicians (non-autoimmune ILDs: 243 pulmonologists; autoimmune ILDs: 243 pulmonologists and 203 rheumatologists). Survey question: "For the following types of ILDs where patients also have lung fibrosis, please indicate your preferred first, second, and third line treatments for the respective ILD". Rheumatologists were only asked this question in relation to RA-ILD and SSc-ILD. Abbreviations. HP, Hypersensitivity pneumonitis; IIP, Idiopathic interstitial pneumonias; ILD, Interstitial lung disease; iNSIP, Idiopathic non-specific interstitial pneumonia; RA, Rheumatoid arthritis; SSc, Systemic sclerosis.

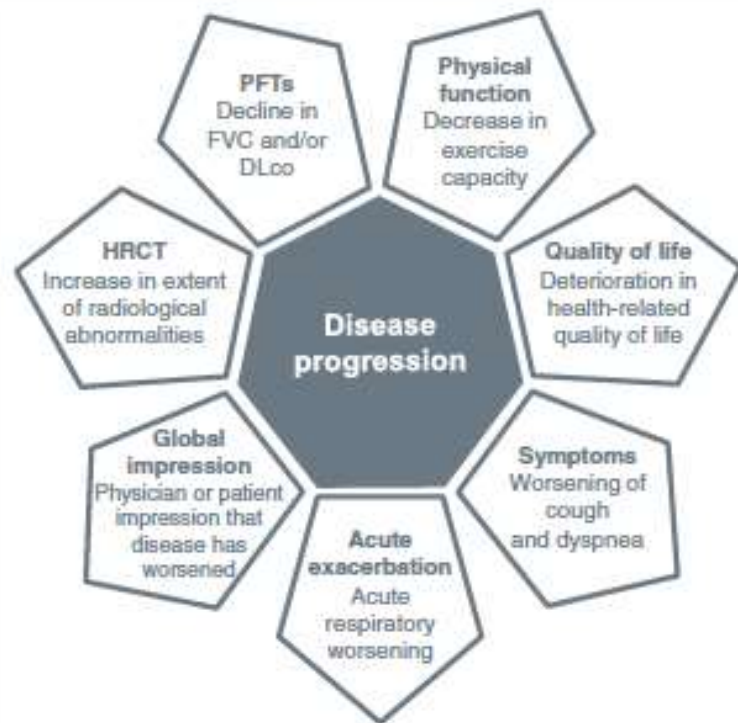
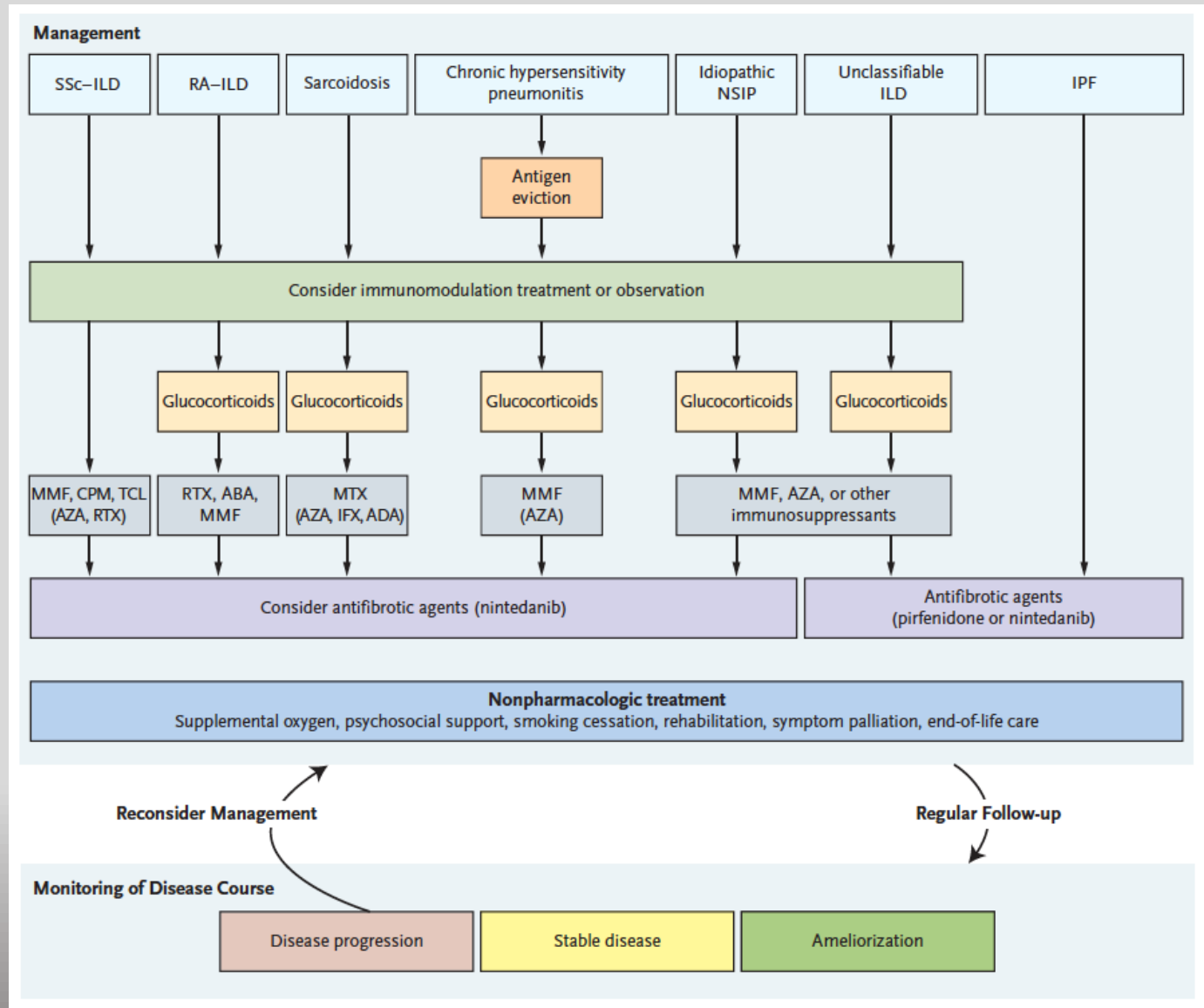


Fig. 2 Factors that reflect progression of interstitial lung diseases. DLco, diffusion capacity of the lungs for carbon monoxide; FVC, forced vital capacity; HRCT, high-resolution computed tomography; PFTs, pulmonary function tests

Kolb Respir Res 2019



Étude INBUILD

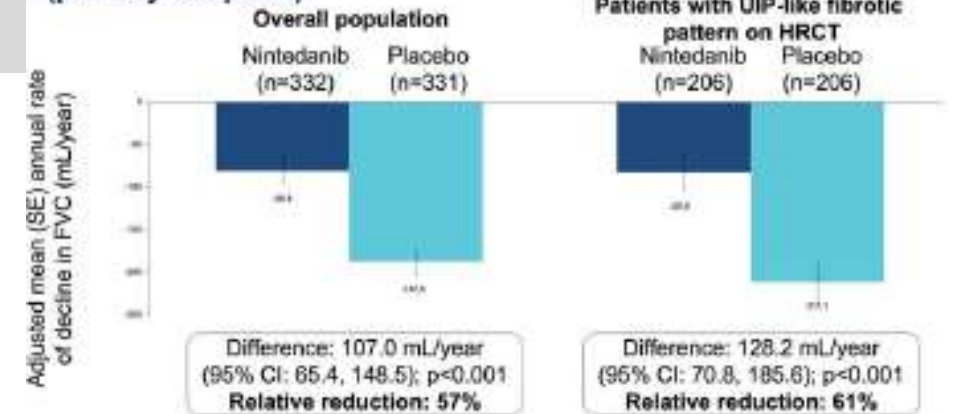
Key inclusion criteria

- Physician-diagnosed ILD other than IPF
- Features of diffuse fibrosing lung disease of >10% extent on HRCT performed ≤12 months prior to screening, confirmed by central review
- FVC ≥45% predicted
- DLco ≥30%–<80% predicted
- Progressive phenotype
 - Patients were required to meet ≥1 of the following criteria for ILD progression in the 24 months before screening, despite management:
 - Relative **decline** in **FVC ≥10%** predicted
 - Relative **decline** in **FVC ≥5–<10%** predicted **AND Worsened** respiratory **symptoms***
 - Relative **decline** in **FVC ≥5–<10%** predicted **AND Increased** extent of **fibrosis on HRCT****
 - **Worsened** respiratory **symptoms*** **AND Increased** extent of **fibrosis on HRCT****

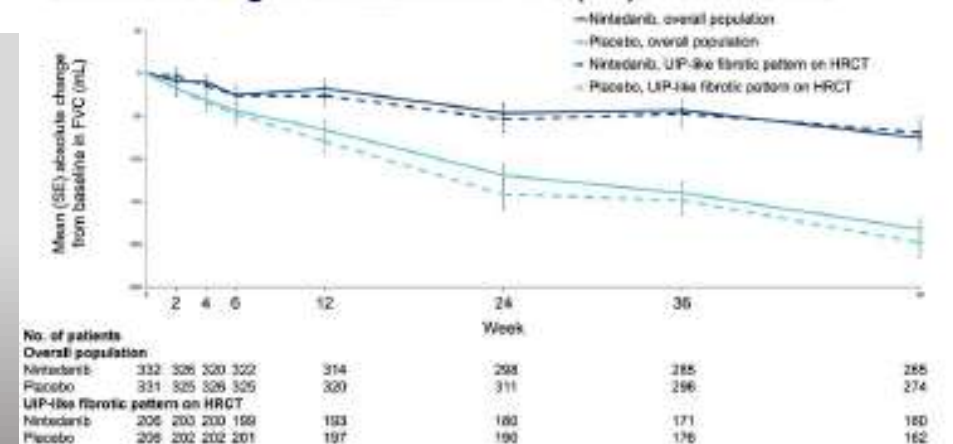
Critères de PID fibrosante

FIBROSING LUNG DISEASE ON HRCT Disease extent of >10%		• Reticular abnormality + Traction bronchiectasis • ± Honeycombing
	Co-existing features allowed	• Ground glass opacity • Upper lung or peribronchovascular predominance • Mosaic attenuation • Air trapping • Centrilobular nodules
	Not allowed	• Widespread consolidation • Progressive massive fibrosis

Adjusted annual rate of decline in FVC (mL/year) over 52 weeks (primary endpoint)



Observed change from baseline in FVC (mL) over 52 weeks



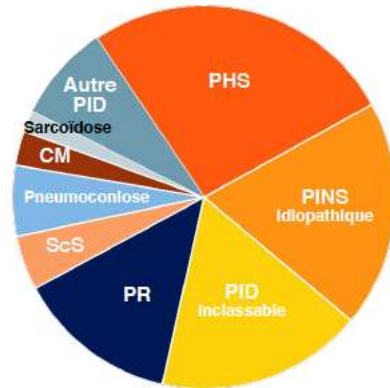
Étude INBUILD

INBUILD Étiologies de la PID à l'inclusion

Nintedanib in patients with progressive fibrosing interstitial lung diseases—subgroup analyses by interstitial lung disease diagnosis in the INBUILD trial: a randomised, double-blind, placebo-controlled, parallel-group trial

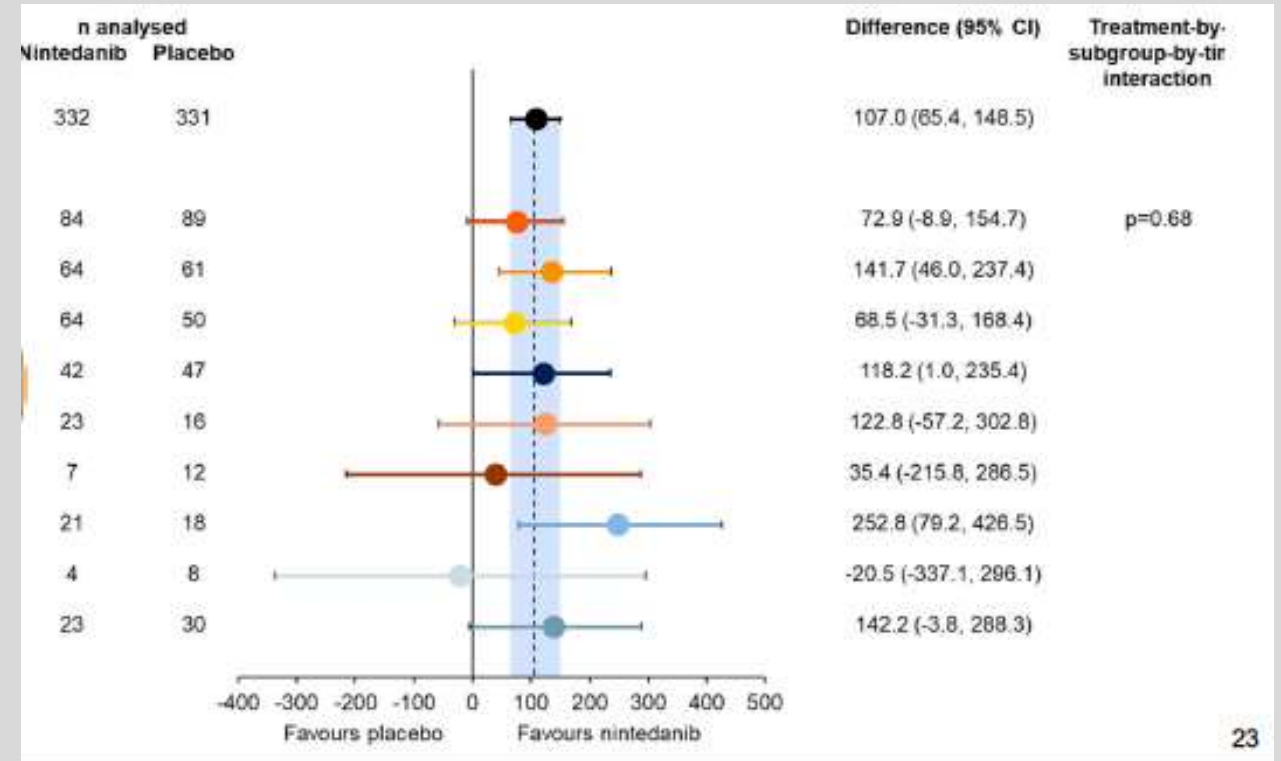
Lancet Respir Med 2020
Published Online
March 5, 2020

Population INBUILD	
PID auto-immune	25%
Associée à une PR	13% (n=42/47)
Associée à une ScS	6% (n=23/16)
Associée à une CTD mixte	3% (n=7/12)
Autre PID auto-immune	3% (n=10/13)
Sarcoidose	2% (n=4/8)
PHS	26% (n=84/89)
Pneumoconiose	6% (n=21/18)
PINS Idiopathique	16% (n=64/50)
PID Inclassable	17% (n=64/50)
Autre PID fibrosante	5% (n=13/17)



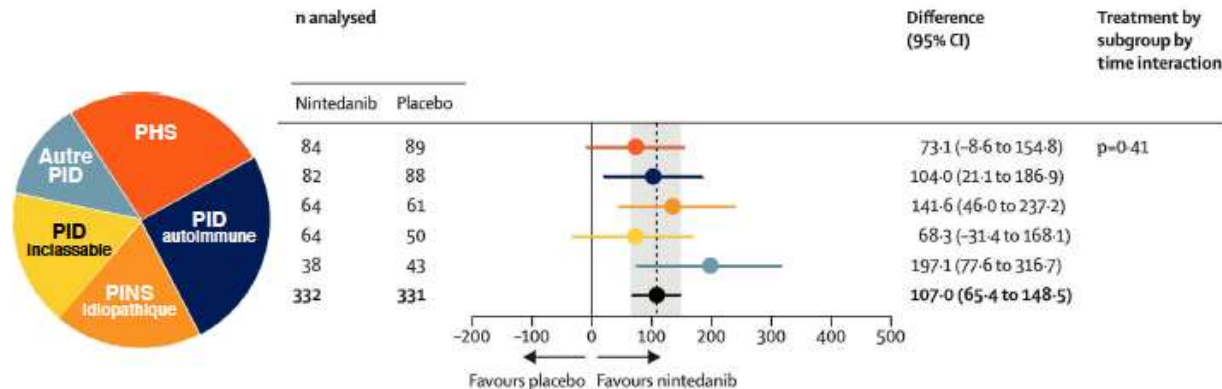
Wells AU, et al. Nintedanib in patients with progressive fibrosing interstitial lung diseases—subgroup analyses by interstitial lung disease diagnosis in the INBUILD trial: a randomised, double-blind, placebo-controlled, parallel-group trial. Lancet Respir Med 2020. [Epub ahead of print]

20



23

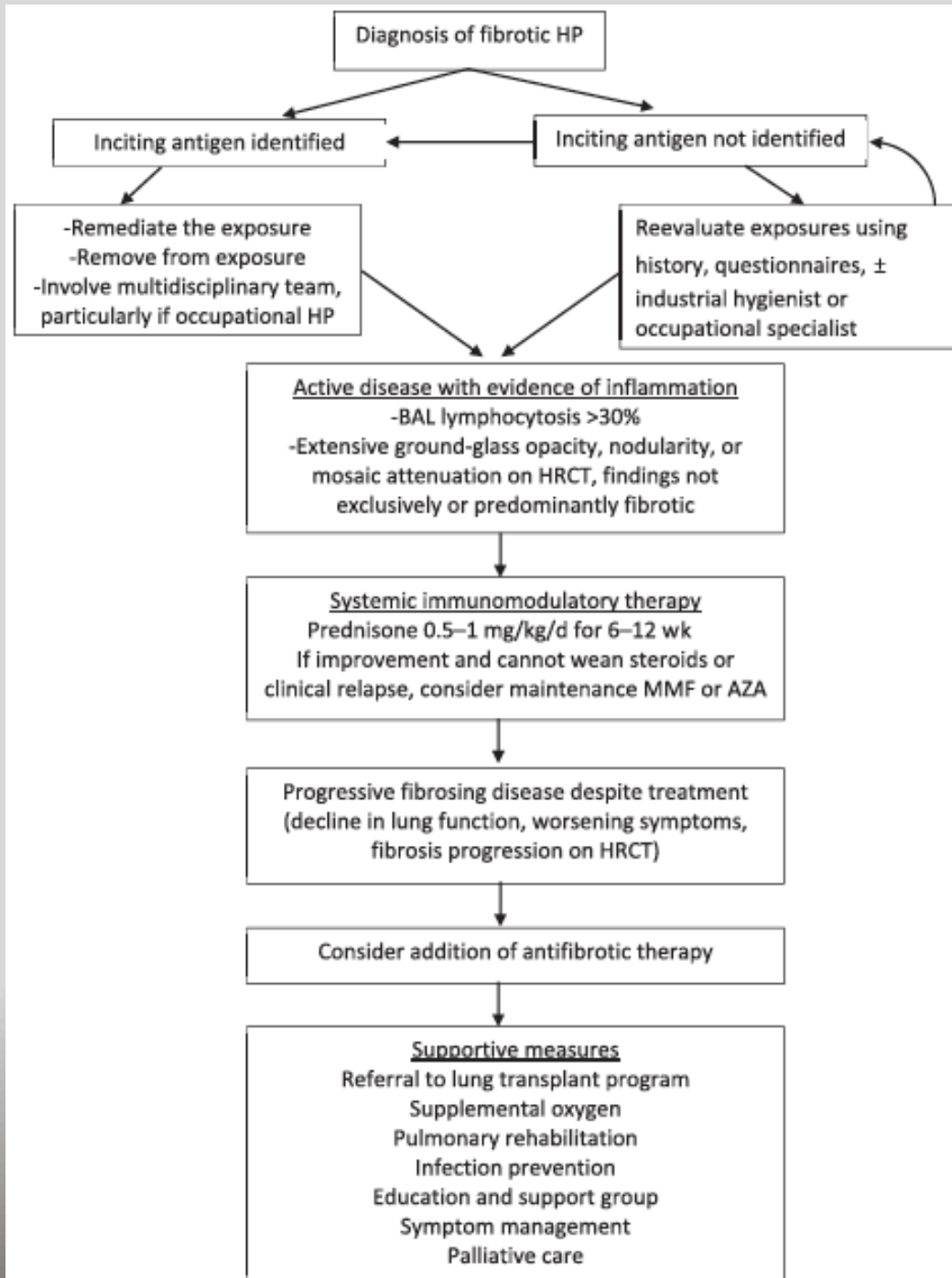
Taux annuel de déclin de la CVF (mL / an)



Management of Fibrotic Hypersensitivity Pneumonitis

Hayley Barnes, MBBS, MPH^{a,b,*}, Kerri A. Johannson, MD, MPH^{c,d}

Clin Chest Med 42 (2021) 311–319



CONCLUSION

Aspect de PINS
et 3 densités ?

Pattern de réticulations mais
péribroncho-vasculaire
Quelle diagnostic FINAL ?

Attention au DD

Topographie des
hypoatténuations

Merci !

Mosaic attenuation
indicatif de SAD:
hypoatténuation

Morphologie lobulaire
Modifications des contrastes
Expiration ?