Population-Based Analysis of Invasive Fungal Infections, France, 2001–2010

Technical Appendix 1: Methods

1. The French Hospital Information System

1a. The Database

The national hospital discharge database PMSI (*Programme de médicalisation des systèmes d’information*) systematically collects information for any new admission: anonymous patient identifier, the hospital code, the patient’s gender, age, residence area, main cause of admission, other medical conditions reported during a stay (>20 entries), acts performed during the stay, duration of stay and mode of discharge including transfer or death when death occurs at hospital. The combination of diseases and acts provides a “discharge summary code” which infers the cost of each stay, provides useful information on major surgery, and allows to check the consistency of collected information (only available from 2004 onwards). This discharge summary code is adapted from the American DRG classification.

The 10th international diseases classification (ICD10) codes are used. These codes were stable over the study period, except for a few variables such as body mass index thresholds for morbid obesity, introduced in 2009.

Medical and surgical acts performed during each stay are coded using the national health insurance classification, which evolved over time. For instance, autologous vs. allogenic hematopoietic stem cell transplant (HSCT) recipients were not distinguished before 2003, and were not systematically differentiated after this period.

1b. Selection of Incident Cases

The patient anonymous identifier, available since 2003, was used to distinguish first admissions from re-hospitalizations. For previous years another identifier was created by chaining variables “year of birth” (derived from the patient’s age and date of admission), “gender” and “residence zip code.” Duplicates were checked within and between hospitals.
Cases detected once over the period were "unique cases", those detected >1 time were "newly admitted case" at first occurrence and “re-admitted” at subsequent occurrence(s).

For re-admitted patients, delays between subsequent stays were estimated from admission and discharge dates. By convention, a case with subsequent stays was defined as a single episode if delay was ≤180 days and as a new episode after this delay. As the proportion of new episodes was very low (<5% of each selected IFI, <8% for Pneumocystis jiroveci pneumonia), we did not retain these new episodes in the final analysis.

1c. Finalisation of the Database with 5 Selected IFIs

The dataset was first created for each selected IFI, including the identification of unique and first admissions (incident cases), the hierarchisation of risk factors and the reporting of death.

In the resulting merged dataset including the five IFIs, when one IFI was reported as principal diagnosis and another one was reported as “associated disease”, priority was given to the first one. When two IFIs were reported as “associated diseases”, priority was given to the rarest one, in order to reduce the under-estimation bias for these rare infections. For instance, a case was considered as mucormycosis when mucormycosis and candidemia were recorded. This process only concerned <1% of the overall dataset.

2. Estimations of Incidence and Trends: Data Sources

• Annual incidence rates by gender and age groups were estimated using the 1999 national population census and its updates, available at the national public health institute (InVS). For neonates (0-28 days) we hypothesised their number was equal to the number of live births.

• Incidence in specific groups was estimated for the following conditions for which annual numbers were available:
  - HIV/AIDS: subject to mandatory notification. Data retrieved from the routine surveillance system, available at InVS (http://www.invs.sante.fr)
Solid tumors patients: we used as a proxy, the number of cancer patients regularly followed up in each hospital. Data available at the Ministry of Health’s division of statistics website (http://www.drees.sante.gouv.fr/statistique-annuelle-des-établissements-sae,6506.html)

Chronic renal failure patients: data provided by National agency in charge: Agence de la Biomédecine (http://www.agence-biomedecine.fr/agence/english.html)

Diabetes: prevalence data on treated diabetic patients and point estimates for all cases obtained from two studies (1,2)

Hematopoietic stem cell transplant (HSCT) recipients: data provided by National agency in charge: Agence de la Biomédecine (http://www.agence-biomedecine.fr/agence/english.html)

3. Case Definitions for Selected IFIs: ICD-10 codes

**Invasive Aspergillosis**: Invasive pulmonary (B44.0) or Disseminated aspergillosis (B44.7).

**Candidemia**: Candidal sepsis (B37.7).

**Candidal meningitis** (B37.5) and endocarditis (B37.6) were excluded from analysis to allow international comparisons.

**Cryptococcosis**: All codes B45: Pulmonary (B45.0), Cerebral incl. meningitis (B45.1), Cutaneous (B45.2), Osseous (B45.3), Disseminated (B45.7), Other (B45.8) or Unspecified forms (B45.9).

**Pneumocystis pneumonia**: Pneumonia due to *Pneumocystis carinii* / P. jiroveci (B59) or HIV disease resulting in *Pneumocystis jirovecii* pneumonia (B20.6) and Confirmed pneumonia

**Zygomycosis/mucormycosis**: All B46 except gastrointestinal (B46.2). All other codes included i.e. Pulmonary (B46.0), Rhinocerebral (B46.1), Cutaneous/subcutaneous (B46.3), Disseminated (generalized) (B46.4) or Unspecified, (B46.5; B46.8; B46.9).

Gastro-intestinal forms were excluded based on the retrospective study ("RetroZygo") among patients diagnosed in metropolitan France from 2005 to 2007 (3,4), in which cases recorded in the PMSI were identified and discussed with the medical teams. The vast majority of gastro-intestinal cases were secondarily excluded as false positives.
4. Case Definitions for Associated Diseases (Risk Factors for IFI)

We used ICD-10 codes, as well as the codes from the National health insurance classification (NHIC) which provided acts performed during a stay (surgical acts, transplantations).

2.1. Hematological Malignancies (HM)


NHIC acts: Chemotherapy for acute leukemias, lymphomas, other tumours or myeloproliferative disorders.

2.1.b. HM with Hematological stem cell transplantation

ICD-10: Bone marrow transplantation, Z94.8 including autologous (Z94.8.00) and allogenic (Z94.8.01) transplantations. Bone-marrow transplant rejection (T86.0).

NHIC acts: Autologous or allogenic stem cell transplantation. Intravenous injection of cellular therapy products.

2.1.c. HM with neutropenia or medullar aplasia

ICD-10: Acquired pure red cell aplasia, D60. Other aplastic anaemia, D61. Agranulocytosis & functional disorders of polymorphonuclear neutrophils, D70; D71.

2.2. Non-hematological Immunosuppressions

2.2.a. HIV-AIDS

2.2.b. Solid organ transplantation/rejection (other than HSCT)

*ICD-10*: Failure and rejection of transplanted organs and tissues, T86. Transplanted organ and tissue status, Z94.

*NHIC*: Heart, liver, lung, pancreas and renal transplantsations (NHIC).

2.2.c. Solid organ tumours

*ICD-10*: Malignant neoplasms, stated or presumed to be primary, of specified sites, except of lymphoid, hematopoietic and related tissue, C00 to C75. Malignant neoplasms of ill-defined, secondary and unspecified sites, C76 to C80.

2.3. Systemic Inflammatory Diseases

*ICD-10*: Crohn disease, K50. Sarcoidosis, D86. Rheumatoid arthritis, M05 (incl. M05.0; M05.1; M05.3; M05.8; M05.9) & M06. Lupus and vasculitis, M05.2; L930; M30; M31; M32

2.4. Diabetes Mellitus


2.5. Chronic Respiratory Diseases

*ICD-10*: Chronic obstructive pulmonary disease (COPB): J44 (incl. J44.0; J44.1; J44.8; J44.9). Asthma, J45. Cystic fibrosis, E84.

2.6. Chronic Renal Failure


*NHIC*: Renal failure with dialysis or dialysis sessions for chronic renal failure (NHIC)

2.7. Acute Renal Failure

*ICD-10*: Acute renal failure, N17. Post-procedural renal failure, N99.0. Renal failure following abortion and ectopic and molar pregnancy, O08.4. Postpartum acute renal failure, O90.4.

*NHIC*: Acute renal failure without dialysis or dialysis session for acute renal failure.
2.8. Morbid Obesity

*ICD-10*: Obesity due to excess calories with BMI [40-49kg/m²], *E66.01*. Obesity due to excess calories, BMI ≥ 50kg/m², *E66.02*. Drug-induced obesity, BMI [40-49kg/m²], *E66.11*. Drug-induced obesity, BMI ≥ 50kg/m², *E66.12*. Extreme obesity with alveolar hypoventilation, BMI [40-49kg/m²], *E66.21*. Extreme obesity with alveolar hypoventilation, BMI ≥ 50 kg/m², *E66.22*. Other obesities, BMI ≥ 40kg/ m², *E66.81; E66.82*.

*NHIC*: surgical treatments for morbid obesities (bypass, other acts).

2.9. Liver Cirrhosis

*ICD-10*: Alcoholic liver disease, *K70.0*. Alcoholic cirrhosis of liver, *K70.3*. Toxic liver disease with fibrosis and cirrhosis of liver, *K71.7*. Fibrosis and cirrhosis of liver, *K74* (incl. *K74.3 – K74.6*).

2.10. Pancreatitis (acute/chronic)

*ICD-10*: Acute pancreatitis, *K85*. Other diseases of pancreas, *K86* (incl. *K86.0-K86.1*).

2.11. Burns

*ICD-10*: Third degree burn: head and neck *T20.3*, trunk *T21.3*, shoulder and upper limbs *T22.3*, wrist and hand *T23.3*, hip and lower limb *T24.3*, ankle and foot *T25.3*. Burns of multiple regions, at least one burn of third degree mentioned, *T29.3*. Burn and corrosion, body region unspecified, third degree *T30.3*. Burns classified according to extent of body surface involved, *T31.1-31.9*.

2.12. Surgery

*Discharge summary codes*: Head-neck-thoracic surgery, Abdomen-urinary tract and gynaecological surgery, and musculo-skeletal surgery.

2.13. Intensive Care

*ICD-10*: Dependence on respirator, *Z99.1*.

*Summary code*: Injury severity score (ISS)>24.

References

