## **PERFECTA** - Intravenous analgesics for post-operative pain: patterns of use in high-risk populations

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General	
Identification	
Detailed name	Intravenous analgesics for post-operative pain: patterns of use in high-risk populations
Sign or acronym	PERFECTA
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	
General Aspects	
Health determinants	Iatrogenic
Others (details)	Pain
Keywords	Post-chirurgical, treatment for pain, IV pain killers
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Unit	Bristol-Myers Squibb
Collaborations	
Funding	
Funding status	Private
Details	Bristol-Myers Squibb
Governance of the database	
Sponsor(s) or organisation(s) responsible	Bristol-Myers Squibb France (BMS)
Organisation status	Private
Additional contact	
Main features	
Type of database	
Type of database	Study databases
Study databases (details)	Not-repeated cross-sectional studies (except case control studies)
Database recruitment is carried out by an intermediary	A selection of health institutions and services
Database recruitment is carried out as part of an interventional study	No
Additional information regarding sample selection.	random sampling in clusters
Database objective	
Main objective	Describe the post-chirurgical drug treatment for pain after a surgical intervention, in target groups of patients having specific risks of developing complications during the use of certain classes of analgesics
Inclusion criteria	M/F patients, >=18 years, have an ASA score of level 3 or 4 and having undergone a surgical

	intervention, including coeloscopy, excluding cardiac surgery and neurosurgery, requiring antalgic treatment through IV in order to relieve postoperative pain. Patients having at least one of the following risk factors: aged ? 65 years, OCPD including sleep apnea, high blood pressure, kidney, heart or liver failure.
Population type	
Age	Adulthood (19 to 24 years) Adulthood (25 to 44 years) Adulthood (45 to 64 years) Elderly (65 to 79 years) Great age (80 years and more)
Population covered	Sick population
Gender	Male Woman
Geography area	National
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Detail of the geography area	Metropolitan France
Detail of the geography area	
Detail of the geography area  Data collection	
Detail of the geography area  Data collection  Dates  Date of first collection (YYYY or	Metropolitan France
Detail of the geography area  Data collection  Dates  Date of first collection (YYYY or MM/YYYY)  Date of last collection (YYYY or	Metropolitan France  2007
Detail of the geography area  Data collection  Dates  Date of first collection (YYYY or MM/YYYY)  Date of last collection (YYYY or MM/YYYY)	Metropolitan France  2007  2008
Detail of the geography area  Data collection  Dates  Date of first collection (YYYY or MM/YYYY)  Date of last collection (YYYY or MM/YYYY)  Size of the database  Size of the database (number of	Metropolitan France  2007  2008
Detail of the geography area  Data collection  Dates  Date of first collection (YYYY or MM/YYYY)  Date of last collection (YYYY or MM/YYYY)  Size of the database  Size of the database (number of individuals)  Details of the number of	Metropolitan France  2007  2008  [1000-10 000[ individuals

Clinical data

Direct physical measures

Type of data collected

Clinical data (detail)

Presence of a biobank	No
Health parameters studied	Health event/morbidity Health event/mortality
Procedures	
Data collection method	paper CRF
Participant monitoring	No
Links to administrative sources	No
Promotion and access	
Promotion and access Promotion	
Promotion	publications
Promotion  Access  Terms of data access (charter for data provision, format of	publications  Access on specific project only