

# Office-Based Surgery - Adverse Event Report

**1. Type of Reportable Adverse Event being reported (please check all that apply):**

- Patient death within 30 days    Date of Death Relative to OBS Procedure within:     24hrs     72 hrs     4-7days     8-30days
- Unplanned transfer to a hospital    Transferring EMS Service: \_\_\_\_\_
- Unscheduled hospital admission for longer than 24 hours w/in 72 hours of undergoing OBS Procedure
- Any serious or life-threatening event (Examples include: permanent or temporary loss of organ or limb function or mental impairment, wrong site surgery and retained foreign body)
- Any suspected transmission of a bloodborne pathogen (BBP) from a health care professional to a patient or between patients. See BBP addendum at the end of this form.

HOSPITAL NAME \_\_\_\_\_

ADDRESS 1 \_\_\_\_\_

ADDRESS 2 \_\_\_\_\_

CITY \_\_\_\_\_

STATE \_\_\_\_\_

ZIP \_\_\_\_\_

**2. Procedure Name(s) and Code(s) Performed on involved patient:**

Procedure Name: \_\_\_\_\_ CPT/HCPCS Code: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

**3. Approximately how many of the type of procedure(s) involved in this report does the primary proceduralist associated with this report perform per month: \_\_\_\_\_**

**4. Date OBS Procedure(s) performed:** \_\_\_\_\_  
MM/DD/YY

**5. Indication for the procedure:**     Screening     Diagnostic     Therapeutic     Optional

**6. Date of Adverse Event:**

Transfer _____ MM/DD/YY	Admission _____ MM/DD/YY
Serious or life threatening event _____ MM/DD/YY	Death _____ MM/DD/YY

**7. Describe events and suspected complication(s) leading up to the unplanned transfer, unscheduled admission, death or serious or life-threatening event reported above. If pregnancy-related procedure, include gestational age of fetus. Attach additional pages if needed.**

**8. Was the (suspected) complication or reason for the adverse event reported (E.g. transfer, admission) identified to the patient as part of the pre-procedure informed consent process?     Yes     No**

**9. Sedation/Anesthesia Related Care**

**a. Significant Past Medical History/Co-morbidity(s):**

- MEDICAL     None     HTN     DM     DVT     Pain     Seizures     Other: \_\_\_\_\_  
 ESRD     CHF     Asthma/COPD     Stroke     Bleeding     Psychiatric    \_\_\_\_\_  
 CAD     Arrhythmias     Anemia     Advanced Stage Cancer     Diverticulitis     Obesity    \_\_\_\_\_  
 PAD     Pregnancy History: G: \_\_\_\_\_ P: \_\_\_\_\_

**b. Patient's Current Home Medications:**

- None     Calcium Channel Blocker     NSAID/ASA     Steroids  
 ACE/ARB     Diuretic     Opiate     Bronchodilators  
 Beta Blocker     Insulin/Oral Hypoglycemic     Anticoagulant     Other: \_\_\_\_\_

**c. Number of hours since last pre-procedure PO intake:**     Less than 6 h     6-12 h     Greater than 12 h

d. ASA Score:  P1  P2  P3  P4  P5  P6  Emergency

e. Pre-procedure Medication(s) Prescribed +/- Administered:  Anxiolytic  Antibiotic  Steroids  
 Antihistamine  Anticoagulant  Other: \_\_\_\_\_

f. Intra and Post Procedural Medications Administered:

1. Procedural sedation/anesthesia medications

DOSE	DOSE	DOSE
_____ Diazepam	_____ Lorazepam	_____ Succinylcholine
_____ Fentanyl	_____ Ketamine	_____ Nitrous Oxide
_____ Morphine	_____ Propofol	_____ "Caine"
_____ Non-depolarizing muscle relaxant	_____ Midazolam	_____ Other _____
_____ Volatile anesthetic agent	_____ Meperidine	

2. Other Medications

DOSE	DOSE	GIVEN
_____ Contrast	_____ Nitroglycerin	<input type="checkbox"/> ACLS/Rescue medication
_____ Flumazinil	_____ Ondansetron/Zofran	<input type="checkbox"/> Antibiotic(s)
_____ Glycopyrrolate/Rubinol	_____ Pitocin/Oxytocin	<input type="checkbox"/> Antihistamine(s)
_____ Heparin	_____ Protamine	<input type="checkbox"/> Bronchodilator(s)
_____ Methergine	_____ tPA	<input type="checkbox"/> Diuretic(s)
_____ Metoclopramide/Reglan	_____ Other _____	<input type="checkbox"/> Steroid(s)
_____ Naloxone/Narcan	_____ Other _____	<input type="checkbox"/> NSAID(s)

g. Level of Anesthesia Achieved:  Local/Regional  Minor  Moderate  Deep  General

10. Length of Procedure:  < 1 hour  1 - 3 hours  3 - 6 hours  > 6 hours

11. Liposuction Volume Removed:  None  <500 ml  501 - 1000 ml  >1000 ml

12. Practitioners participating in reported OBS procedure:

a. Name of Primary proceduralist/surgeon:

\_\_\_\_\_  
LAST, FIRST, MI LICENSE/CERT. TYPE LICENSE #

b. Practitioner who determined anesthetic/sedative drug(s) and dosages:

\_\_\_\_\_  
LAST, FIRST, MI LICENSE/CERT. TYPE LICENSE #  
Credential/Role:  BC/BE/Anesthesiologist  Proceduralist  CRNA  Other: \_\_\_\_\_

c. Practitioner who administered anesthesia/sedation:

\_\_\_\_\_  
LAST, FIRST, MI LICENSE/CERT. TYPE LICENSE #

d. Others participating in procedure:

\_\_\_\_\_  
LAST, FIRST, MI CERT/TITLE LICENSE #

\_\_\_\_\_  
LAST, FIRST, MI CERT/TITLE LICENSE #

\_\_\_\_\_  
LAST, FIRST, MI CERT/TITLE LICENSE #

13. Patient Information:

\_\_\_\_\_  
LAST, FIRST, MI GENDER AGE DOB (DD/MM/YY) LAST 4 SSN DIGITS

\_\_\_\_\_  
ADDRESS

\_\_\_\_\_  
CITY STATE ZIP

**14. Location where OBS was performed:**

LEGAL NAME \_\_\_\_\_ PRACTICE NPI # \_\_\_\_\_  
ADDRESS 1 \_\_\_\_\_  
ADDRESS 2 \_\_\_\_\_  
CITY \_\_\_\_\_ STATE \_\_\_\_\_ ZIP \_\_\_\_\_  
Name of Contact Person:  
LAST, FIRST, MI \_\_\_\_\_ PHONE W/AREA CODE \_\_\_\_\_ EMAIL \_\_\_\_\_

**15. Quality Improvement:**

- a. In the opinion of the proceduralist/surgeon, or reporter if this report is being filed by a practitioner or facility not affiliated with the practice where the OBS occurred, the adverse event being reported is related to the:
- Procedure     Anesthesia/Sedation     Equipment Factors     Patient Factors     Practitioner Factors     System/Practice Factors  
 Other, please identify: \_\_\_\_\_

- b. Please identify actions the Practice/Practitioner has taken/plans to take to prevent similar adverse events from happening in the future:  
\_\_\_\_\_  
\_\_\_\_\_

**16. Accreditation: Effective July 14, 2009 all practices in which office-based surgery is performed must be accredited by an agency designated by the Commissioner of Health.**

- a. Is your practice accredited?     No     Yes, with:     AAAASF     AAAHC     TJC  
b. If not yet accredited, has your practice applied for accreditation?     No     Yes, with:     AAAASF     AAAHC     TJC  
When do you expect to receive your OBS accreditation? \_\_\_\_\_  
DATE

*American Association for the Accreditation of Ambulatory Surgical Facilities (AAAASF); Accreditation Association for Ambulatory Health Care (AAAHC); The Joint Commission (TJC)*

**17. Reporter(s): Name(s) and Signature(s) of Reporters Attesting to the Accuracy of this Report:**

*(All MD, PA, SA that participated in the procedure must report; each practitioner can submit a report or multiple practitioners can sign the same report.)*

PRINTED NAME \_\_\_\_\_ SIGNATURE \_\_\_\_\_  
Practice/Facility Affiliation:  
 Practice where OBS occurred     Other \_\_\_\_\_  
PRACTICE NAME \_\_\_\_\_  
STREET \_\_\_\_\_  
CITY, STATE, ZIP \_\_\_\_\_ PHONE \_\_\_\_\_

PRINTED NAME \_\_\_\_\_ SIGNATURE \_\_\_\_\_  
Practice/Facility Affiliation:  
 Practice where OBS occurred     Other \_\_\_\_\_  
PRACTICE NAME \_\_\_\_\_  
STREET \_\_\_\_\_  
CITY, STATE, ZIP \_\_\_\_\_ PHONE \_\_\_\_\_

*Item 17 continued on next page.*

PRINTED NAME

SIGNATURE

Practice/Facility Affiliation:

Practice where OBS occurred

Other

PRACTICE NAME

STREET

CITY, STATE, ZIP

PHONE

PRINTED NAME

SIGNATURE

Practice/Facility Affiliation:

Practice where OBS occurred

Other

PRACTICE NAME

STREET

CITY, STATE, ZIP

PHONE

**18. Date of Report:**

DD/MM/YY

*Adverse event reports must be reported w/in 24 hours of the event. If this report is filed more than one business day after the event, provide a description of the factors that prevented you from filing the report within the required timeframe.*

Reason for delay in reporting, if applicable:

Please submit signed Adverse Event form, via certified mail, to:

New York State Department of Health  
Patient Safety Center  
Hedley Park Place  
433 River St  
Troy, NY, 12180

**ADDENDUM: Bloodborne Pathogen Transmission**

**Date of Suspected Transmission of Bloodborne Pathogen:**

MM/DD/YY

**Full Description of Events related to Suspected Transmission of BBP (attach additional pages if needed):**

**Bloodborne pathogen involved in (suspected) transmission:**

**Type of Transmission:**

Healthcare professional to patient

Between or among patients

**Number of Patients Affected:**