

**PART I - ESTABLISHMENT INFORMATION**

3. OTHER FDA REGISTRATIONS

a. BLOOD FDA 2830 NO. \_\_\_\_\_

b. DEVICES FDA 2891 NO. \_\_\_\_\_

c. DRUG FDA 2656 NO. \_\_\_\_\_

4. PHYSICAL LOCATION (Include legal name, number and street, city, state, country, and post office code)  
 Sighthlife (also dba Sighthlife Surgical Inc.)  
 6000 Shoreline Ct Suite 202  
 South San Francisco, California 94080

a. PHONE 650-276-7900 EXT \_\_\_\_\_

b.  SATELLITE RECOVERY ESTABLISHMENT  
 (MANUFACTURING ESTABLISHMENT FEI NO. \_\_\_\_\_)

c.  TESTING FOR MICRO-ORGANISMS ONLY

5. ENTER CORRECTIONS TO ITEM 4

6. MAILING ADDRESS OF REPORTING OFFICIAL (Include institution name if applicable, number and street, city, state, country, and post office code)  
 Sighthlife  
 Attn: Thomas D. Miller, B.S. CBET  
 1200 6th Ave  
 Ste 300  
 Seattle, Washington 98101


7. ENTER CORRECTIONS TO ITEM 6

a. PHONE 206-838-4630 EXT \_\_\_\_\_

b. PHONE \_\_\_\_\_

8. U.S. AGENT

a. E-MAIL \_\_\_\_\_

9. REPORTING OFFICIAL'S SIGNATURE  
  
 a. TYPED NAME Thomas D. Miller, B.S. CBET  
 b. E-MAIL tom.miller@sighthlife.org  
 c. TITLE VP of Quality and Regulatory Affairs  
 d. DATE 21-DEC-2016

**PART II - PRODUCT INFORMATION**

Types of HCT / Ps	Establishment Functions					11 HCT/PS DESCRIBED IN 21 CFR 1271.10	12 HCT/PS REGULATED AS MEDICAL DEVICES	13 HCT/PS REGULATED AS BIOLOGICAL DRUGS	14. PROPRIETARY NAME(S)
	Recover	Screen	Test	Package	Process				
a. Bone									
b. Cartilage									
c. Cornea	X	X		X	X			X	
d. Dura Mater									
e. Embryo									
f. Fascia									
g. Heart Valve									
h. Ligament									
i. Oocyte									
j. Pericardium									
k. Peripheral Blood Stem									
l. Sclera	X	X		X	X			X	
m. Semen									
n. Skin									
o. Somatic Cell Therapy Products									
p. Tendon									
q. Umbilical Cord Blood									
r. Vascular Graft									
s. Amniotic Membrane									AmnioGraft, PROKERA
t.									
u.									
v.									