

የናርኮቲክ መድኃኒቶችንና የሳይኮትሮፒክ ንጥረ ነገሮችን ለመቆጣጠርና በአግባቡ ጥቅም ሳይ ለማዋል የወጣ መመርያ 370/2013

DIRECTIVES TO CONTROL AND PROMOTE PROPER USE OF NARCOTIC DRUGS AND PSYCHOTROPIC SUBSTANCES

የኢትየጵያ የምግብ፣ የመድሃኒትና የጤና ክብካቤ አስተዳደርና ቁጥጥር ባለስልጣን

FOOD, MEDICINE AND HEALTHCARE ADMINISTRATION AND CONTROL AUTHORITY OF ETHIOPIA

አዲስ አበባ Addis Ababa ጥቅምት 2006 October, 2013

መግቢያ

የናርኮቲክ መድኃኒቶችንና የሣይኮትሮፒክ ንዋረ ነገሮችን ሕገወዋ ምርት፣ ክፍፍልና አጠቃቀም መከላከልና መቆጣጠር አስፈላጊ ሆኖ በመገኘቱ፣

የናርኮቲክ መድኃኒቶችና የግይኮትሮፒክ ንዋረ ነገሮችን ምርት አላሳክ፣ አከፋሬል፣ አስተዛዘዝ፣ ዕደላና አጠቃቀም ስርዓት ባለው መንገድ እንዲከናወን በማስፈለጉ፣

ለዚህም አስተማማኝ የሆነ የናርኮቲክ መድኃኒቶችና የሣይኮትሮፒክ ንዋረነገሮችን ቁዋዋር ስርዓት መዘርጋት አስፈላጊ ሆኖ በመገኘቱ፣

በምግብ፣ የመድኃኒትና ጤና ክብካቤ አስተዳደር ቁጥጥር አዋጅ ቁጥር 661/2002 አንቀጽ 55(3) መሰረት ይህ መመሪያ ወጥቷል፡፡

ክፍል አንድ

ጠቅሳሳ

1. አጭር ርዕስ

ይህ መመሪያ "የናርኮቲክ መድኃኒቶችንና የግይኮትሮፒክ ንጥረ ነገሮችን ለመቆጣጠርና በአግባቡ ጥቅም ሳይ ለማዋል የወጣ መመሪያ ቁጥር 370/2013 ተብሎ ሊጠቀስ ይችሳል።

2. **ትርጓሜ**፣

የቃሉ አገባብ ሌላ ትርጉም የሚያሰጠው ካልሆነ በስተቀር በዚህ መመሪያ ውስጥ፤

- 1. "የናርኮቲክ መድኃኒት" ማለት ኢትዮጵያ በተቀበለችው የተባበሩት መንግስታት የናርኮቲክ መድኃኒቶች ቁጥጥር ስምምነት መሰረት አለም አቀፍ ቁጥጥር የሚደረግበት መድኃኒት ሲሆን ባለስልጣኑ የናርኮቲክ መድኃኒት ብሎ የሚሰይመውንም ይጨምራል።
- 2. "የሣይኮትሮፒክ ንዋረ ነገር" ማለት ኢትዮጵያ በተቀበለችው የተባበሩት መንግስታት የሣይኮትሮፒክ ንዋረ ነገሮችን ቁዋዋር ስምምነት መሰረት አለም አቀፍ ቁዋዋር የሚደረግበት መድኃኒት ሲሆን ባለስልጣኑ የሣይኮትሮፒክ ንዋረ ነገር ብሎ የሚሰይመውንም ይጨምራል፡፡
- 3. "የጤና ተቋም" ማለት የጤና ማበልጸግ፣ የበሽታ መከላከል፣ ማከምና መልሶ ማቋቋም ስራዎችን ወይም የመድኃኒት ንግድ ስራን ወይም አገልግሎት የሚያከናውን ማንኛውም የመንግስት፣ መንግስታዊ ያልሆነ ወይም የግል ተቋም ነው፡፡
- 5. " አግባብ ያለው አካል" ማለት እንደአግባቡ አስፌጻሚ አካሉ ወይም የምግብ፣ መድ ኃኒትና የጤና ነክ ቁጥጥር የሚደረግበት ተቋም ተግባራትን በክልል ደረጃ ማከናወን ሥልጣን የተሰጠው የክልል መንግሥትአካል ወይም በህግ ሥልጣን የተሰጠው ሌላ አካል ነው፡፡
- 6. "የመድኃኒት ንግድ ተቋም" ማለት የናርኮቲክ መድኃኒቶች ወይም ሳይኮትሮፒክ ንዋረ

7. ነገሮች ለማስመጣት፣ ለመሳክ፣ ለማምረት፣ ለማከፋፈል፣ ለማከማቸት፣ ለመያዝ ወይም ለመቸርቸር ከሚመለከተው አካል የብቃት ማረጋገጫና የንግድ ፍቃድ የተሰጠው ተቋም ነው።

ክፍል ሁለት

ስለ ልዩ ፌቃድና የፌቃድ አሰጣጥ

3. የልዩ ፌቃድ አስፈላጊነት

- 1) ማንኛውም ሰው ናርስቲክ መድኃኒቶች ወይም ሳይስትሮፒክ ንዋረ ነገሮች ለማስመጣት፣ ለመሳክ፣ ለማምረት፣ ለማከፋፌል፣ ለማከማቾት፣ ለመያዝ፣ ለኬሚካል ምርመራ ወይም ለዋናታዊ ምርምር ለመጠቀም ከባለስልጣኑ ልዩ ፌቃድ ማውጣት አለበት፡፡
- 2) በዚህ አንቀጽ ንዑስ አንቀጽ 1 መሥረት ልዩ ፌቃድ የሚሰጠው መድኃኒት ማስመጣትን፣ መሳክን፣ ማምረትን፣ ማከፋፌልን ወይም ማከማቸትን ወይም የጤና አገልግሎት መስጠትን በሚመለከት የብቃት ማረጋገጫ የምስክር ወረቀት ላለው ሰው ብቻ ይሆናል፡፡
- 3) ማንኛውም በዚህ አንቀፅ ንዑስ አንቀፅ 1 ስር የተጠቀሱትን ተግባራት ለማከናወን የሚፌልግ ሰው በዚህ መመሪያ አንቀፅ 5 መሰረት ማመልከቻ ማቅረብ አለበት፡፡

4. ልዩ ፌቃድ የሚያስከለክሉ ተግባራት

ማንኛውም ሰው ከዚህ በታች የተዘረዘሩትን ተግባራት ፌፅሞ ሲገኝ የናርኮቲክ መድኃኒትንና የሳይኮትሮፒክ ንዋረ ነገርን ለመያዝ የሚስችለውን ልዩ ፌቃድ ማግኘት አይችልም፡፡

- 1) በሱሰኘንት፣ በሕገወጥ የናርኮቲክ መድኃኒቶች ወይም ሣይኮትሮፒክ ንጥረ ነገሮችን አዘዋዋሪነት፣ ወይም እንዚህኑ ወንጀሎች ለመፈጸም ሙከራ ያደረገ ወይም ያምበረበረ ወይም የተምበረበሩ ማስረጃዎችን የተጠቀመ ወይም ለመጠቀም የሞከረ፣
- 2) የናርኮቲክ መድኃኒቶችን ወይም ሣይኮትሮፒክ ንዋረ ነገሮችን ለመያዝ አመቺ የሆነ ክፍል ወይም ቁሣቁስ ከሌለ፣

5. የማስመጣት ወይም የመሳክ ልዩ ፌቃድ ለማግኘት የሚቀርብ ማመልከቻ

- 1) በዚህ መመሪያ አንቀጽ 3 የተገለጸውን ልዩ ፍቃድ ለማግኘት ለባለስልጣኑ የሚቀርብ ማመልከቻ በሁለት ቅጂ ተዘጋጅቶ የሚቀርብ ሲሆን የሚከተሉት መረጃዎች መያዝ ይኖርበታል፡-
 - (U) የአመልካቹ ሙሉ ስም፣ ሙያ፣ ዜግንት፣
 - (ሰ) የድርጅቱን ስምና አድራሻ፣
 - (ሐ) የንግድ አይነት፣
 - (መ) የመድኃኒቱ ወይም የንተረ ነገሩ ወይም ተሬ ዕቃ ስም፣ ዓይነትና ተንካሬ ማካተት አለበት
- 2) ባለስልጣኑ እንደ አግባቡ የቀረበውን ማመልከቻ ተስተካክሎ በድጋሚ እንዲቀርብ ለመቀነስ ወይም ክብደት ለማሻሻል ወይም አስፈላጊ ያልሆኑትን ከዝርዝር የማውጣት ሥልጣንና ኃላፊነት አለው።

6. የማስመጣት ወይም የመሳክ ልዩ ፌቃድ ስለመስጠት፣

- 1) ባለስልጣኑ ማንኛውም ናርኮቲክ መድኃኒት ወይም ሳይኮትሮፒክ ንዋረ ነገር ወይም ዋሬ ዕቃ ወደ አገር ውስዋ እንዲገባ ልዩ ፍቃድ የሚሰጠው፣
 - (U) መድኃኒቱ በብሔራዊ የመድኃኒት መዘርዝር ውስጥ የተካተተ ከሆነ፣
 - (ለ) መድኃኒቱ በአገሪቱ ውስጥ ለሕክምና አገልግሎት፣ ለሣይንሣዊ ምርምር

ወይም ለሌላ ሕጋዊ ተግባር አስፈላጊ መሆኑ ሲረጋገጥ

- 2) ባለስልጣት ማንኛውም በአገር ውስጥ የተመረተ ናርኮቲክ መድኃኒት ወይም ሳይኮትሮፒክ ንጥረ ነገር ወይም ጥሬ ዕቃ ከአገር እንዲወጣ ልዩ ፌቃድ የሚሰጠው፣ ሳኪው ድርጅት ከተቀባይ አገር ከሚገኘው ተቆጣጣሪ ባለስልጣን ልዩ የማስመጣት ፍቃድ ሲያቀርብ ይሆናል።
- 3) ማመልከቻው ተሟልቶ ከቀረበ በሁለት የስራ ቀናት ውስጥ ድርጅቱ ፌቃድ ያገኛል፣

7. ወደ ሀገር ውስጥ ስለማስገባት

- 1) ማንኛውም ሰው ናርኮቲክ መድኃኒት ወይም ሳይኮትሮፒክ ንዋረ ነገር ወደ ሀገር ውስጥ ለማስገባት:-
 - (ሀ) በባለስልጣኑ የተሰጠ ልዩ የማስገቢያ ፌቃድ፤
 - (ለ) መድኃኒቱ የሚመጣው በአየር መጓጓዣ ሆኖ ለብቻው የታሸባ፣እና
 - (ሐ) ለዚህ አላማ ብቻ የተዘ*ጋ*ጀ ኢንቮይስ፤ ማሟላት አለበት።
- 2) ማንኛውም ሰው ናርኮቲክ መድኃኒት ወይም ሳይኮትሮፒክ ንዋረ ነገር ከመውጫና መግቢያ በር ሳይ ለማስለቀቅ የሚከተሉትን ሰንዶች ከማመልከቻው ጋር አያይዞ ማቅረብ አለበት፤
 - (ሀ) የምዝገባ ምስክር ወረቀት (Registration certeficate) ፤
 - (ለ) ለአያንዳንዱ መለያ ቁጥር የተሰጠው የይዘት ምስክር ወረቀት (Batch Analysis Certificate) ዋና ወይም ቅጅውን ፤
 - (ሐ) የስሪት አገር ማረ,ጋገጫ ሰርተፊኬት (Certeficate of Origin) ፤
 - (መ) የችቃ ዝርዝር መግለጫ ሰንድ (Packing List)፤

- (ሥ) ቢል ኦፍ ሎዲንግ ወይም ኤርዌይ ቢል (Bill of Loading or Airway Bill) ፤
- (ረ) ኢንቮይስ (Comercial Invoice) ፤ እና
- (ሰ) የቅድሙ መግቢያ ፌቃድ ሰርተፊኬት (Pre-import permit certificate)
- 3) ባለስልጣኑ በመውጫና መግቢያ በር ላይ የቀረበለትን ሰነድ ትክክለኛነት በግረ,ጋገጥ የመልቀቂያ ፌቃድ ይሰጣል፡፡ የመልቀቂያ ፌቃድ ኮፒም ለገቢዎችና የጉምሩክ ባለስልጣን እና ለአስመጪው ወይም ለላኪው ድርጅት ይሰጣል፡፡

8. የማስገባት ልዩ ፌቃድ ኮፒዎች ስርጭት

ባለስልጣኑ ናርኮቲክ መድኃኒት ወይም ሳይኮትሮፒክ ንዋረ ነገር ለማስገባት የሚሰጠው ልዩ ፍቃድ በአራት ቅጂ የሚዘጋጅ ሆኖ፣

- 1) ዋናው ቅጂ ለአስመጪው የሚሰጥ ሆኖ አስመጪው ቅጅውን ለሳኪው ድርጅት መሳክ አለበት፡፡
- 2) ባለስልጣኑ ሁለተኛውን ቅጂ ሳኪው ድርጅት ለሚገኝበት አገር ተቆጣጣሪ አካል ይልካል፡፡
- 3) ባለስልጣኑ ሶስተኛውን ቅጂ መድኃኒቱ ወይም ንዋረ ነገሩ ወይም ዋሬ ዕቃው በሚገባበት የመውጫና መግቢያ በር ለሚገኝ የጉምሩክ ባለስልጣን ለክትትል ይላካል፡፡
- 4) አራተኛ ቅጂ ለባለስልጣኑ ቀሪ ይሆናል፡፡

9. ስለ ልዩ ፌቃድ የአገልግሎት ጊዜ ገደብ

1) የናርኮቲክ መድኃኒቶች ወይም የሳይኮትሮፒክ ንዋረ ነገሮችን ለማስመጣት ወይም ለመሳክ የሚሰዋ ልዩ ፌቃድ የሚያገለግለው ከተሰጠበት ቀን ጀምሮ በዘጠና ቀናት ውስዋ አንድ ጊዜ ለማስመጣት ወይም ለመሳክ ብቻ ይሆናል፡፡ 2) የተሰጠው ልዩ ፌቃድ በተጠቀሰው የጊዜ ገደብ ውስጥ ጥቅም ላይ ሳይውል ሲቀርና ለባለስልጣኑ ሲመለስ ወይም የተሰጠው ልዩ ፌቃድ የጠፋ ከሆነ መጥፋቱን የሚያረጋግጥ መረጃ ሲቀርብ የተሰጠው ልዩ ፌቃድ ተሰርዞ በምትኩ አዲስ ልዩ ፌቃድ ሊሰጥ ይችላል፡፡

ክፍል ሶስት

ስለማዘዝና ስለማደል

10. የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንጥረ ነገርን ማደልና ማዘዝ

- 1) ማንኛውም የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋረ ነገርን እንዲያዝ የተፈቀደለት የጤና ባለሙያ መድኃኒትን ወይም ንዋረ ነገርን ማዘዝ የሚችለው ለዚሁ ተብሎ በተዘጋጀው ልዩ የማዘዣ ወረቀት ብቻ ነው፡፡
- 2) ማንኛውም የጤና ተቋም የናርኮቲክ *መድኃኒት*ና የሣይኮትሮፒክ ንዋረ ነገርን ማደል የሚችለው ለዚሁ ተብሎ በተዘ*ጋ*ጀው ልዩ የማዘዣ ወረቀት ብቻ ይሆናል፡፡
- 3) ማንኛውም የጤና ባለሙያ የናርኮቲክ *መድኃ*ኒትና የሣይኮትሮፒክ ንዋሪ ነገር ማዘዣ ወረቀት የተጨበረበረ፣ የተደለዘ ወይም ጊዜው ያለፌበት መሆኑን እያወቀ ማደል ወይም መሸጥ አይችልም፡፡
- 4) ማንኛውም የጤና ባለሙያ የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዯረ ነገርን በተፈቀደለት የሕክምና ባለሙያ ሳልታዘዘለት ሰው ማደል ወይም መሸዋ ወይም እንዲጠቀም ማድረግ የተከለከለ ነው፡፡
- 5) ማንኛውም የጤና ባለሙያ የናርኮቲክ *መድኃ*ኒትና የሣይኮትሮፒክ ንዋሪ ነገር ለራሱ ማዘዝ አይችልም ፡፡

6) ማንኛውም የጤና ባለሙያ የናርኮቲክ *መ*ድኃኒትና የሣይኮትሮፒክ ን**ተ**ረ ነገርለማዘዝ የሚያስችል ልዩ ፌቃድ ቢኖረውም ያለበቂ ምክንያት ወይም ከተገቢው መጠን በላይ ማዘዝ አይችልም ፡፡

11. <u>የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋሪ ነገር ለማዘዝ የተፌቀደለት የጤና</u> ባለሙያ

የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንጥረ ነገር ማዘዝ የሚችለው በሙያው እንዲሰራ ሕጋዊ ፌቃድ ከሚመለከተው አካል ያገኘና የሙያው ደረጃ የሚፌቅድለት የሕክምና ባለሙያ ብቻ ይሆናል፡፡

12. የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንጥረ ነገር አስተዛዘዝ ስርዓት

- 1) ማንኛውም የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋረ ነገር ልዩ ማዘዣ ወረቀት የሚከተሉትን መረጃዎች መያዝ አለበት፣
 - (ሀ) የታዘዘበት ቀን፣
 - (ሰ) የታካሚውን ስም፣ ዕድሜ፣ ጸታ፣ አድራሻና የካርድ ቁጥር፣ የህመሙ ዓይነት ወይም አለም አቀፍ መለያ ኮድ ቁጥር፣
 - (ሐ) የመድኃኒቱ ስም፣ ጥንካሬ፣ የዝግጅት ዓይነት፣ ብዛት፣ የታዘዘለት ቀን ብዛትና የአወሳሰድ መመሪያ በግልጽ የተጻፈበት፣
 - (መ) ያዘዘው ባለሙያ ስም፣ አድራሻ፣ የሙያ ምዝገባ ፍቃድ ቁጥርና ፊርጣ፣
 - (**w**) ያደለው ባለሙያ ስም፣ አድራሻና ፊርጣ፣
- 2) የማዘዣ ወረቀቱ በብዕር የተጻፈና የተፈረመ መሆን አለበት፡፡
- 13. የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንጥረ ነገር ስለማደል

- 1) ማንኛውም የጤና ባለሙያ ልዩ ማዘዣ ወረቀቱ በዚህ መመሪያ አንቀጽ 12 ንዑስ ቁጥር 1 ስር የተጠቀሱትን ካለሟላ በስተቀር የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንጥረ ነገርን ማደል አይችልም፡፡
- 2) ማንኛውም የናርኮቲክ መድኃኒትና የግይኮትሮፒክ ንዋሪ ነገር የታዘዘበት የማዘገና ወረቀት የአገልግሎት ጊዜው ከታዘዘበት ቀን ጀምሮ ለአሥራ አምስት /15/ ቀን ብቻ ነው፡፡
- 3) ማንኛውም የጤና ባለሙያ ዕድሜው ከአሥራ አምስት /15/ ዓመት በታች ለሆነ ሰው ለራሱም ሆነ" ለቤተሰቡ ወይም ለሌላ ሰው የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንጥረ ነገርን መስጠት ወይም ማደል የለበትም፡፡

ክፍል አራት

ስለ ሪኮርድና ሪፖርት

14. ጠቅሳሳ

- 1) ማንኛውም የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋሪ ነገር ያመረተ፣ ያስመጣ፣ የሳከ፣ ወይም ያከፋፈለ ሰው& የየዕለት የሥራ ክንውን መረጃ መያዝ አለበት፡፡
- 2) ማንኛውም ፌቃድ የተሰጠው ሰው ስላመረተው፣ ስላስመጣው፣ ስለላከው ወይም ስላከፋፌለው የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋረ ነገር ወይም ጥሬ ዕቃዎችን ለባለስልጣኑ ሪፖርት ማቅረብ አለበት፡፡

15. ሪኮርድ አያያዝ

- 1) ማንኛውም የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋረ ነገር የመድኃኒት ንግድ ተkም ፣
 - (ሀ) ስላስመጣው ናርኮቲክ መድኃኒት በቅጽ NPS/01/A፣

- (ሰ) ስላስመጣው ሣይኮትሮፒክ ንጥረ ነገር በቅጽ NPS/01/B፣
- (ሐ) ስለሳከው የናርኮቲክ መድኃኒት በቅጽ NPS/02/A፣
- (መ) ስለሳከው ግይኮትሮፒክ ንጥረ ነገር በቅጽ NPS/02/B፣
- (**ሠ**) ስላከፋፈለው የናርኮቲክ መድኃኒት በቅጽ NPS/03/A፣
- (ሬ) ስላክፋፌለው ሣይኮትሮፒክ ንጥሬ ነገር በቅጽ NPS/03/B፣ መረጃ መያዝ አለበት
- 2) ማንኛውም የናርኮቲክ መድኃኒትና የሣይኮትሮፒክ ንዋረ ነገር አምራች ድርጅት፣
 - (ሀ) ያስመጣውንና የተጠቀመውን የናርኮቲክና ግይኮትሮፒክ ጥሬ ዕቃዎች በቅጽ NPS/02/D
 - (ሰ) ያመረተውን ናርኮቲክ መድኃኒት በቅጽ NPS/01/D
 - (ሐ) *ያመረተውን* ሳይኮትሮፒክ ንጥር ነገር በቅጽ NPS/01/E
 - (ሥ) ያስፋፌለውን ናርኮቲክ መድኃኒት በቅጽ NPS/03/A፣ መያዝ አለበት

3) ማንኛውም የጤና ተቋም

- (ሀ) የገዛውን ወይም ያገኘውን የናርኮቲክ መድኃኒት ወይም የሣይኮትሮፒክ ንጥረነገር በሞዴል 19 ወይም በተቋሙ ሕጋዊ የገቢ ደረሰኝ ገቢ ማድረግ አለበት፡፡
- (ለ) የመድኃኒት ማደያ ክፍል ያደለውን የናርኮቲክ መድኃኒት ወይም የሣይኮትሮፒክ ንጥረ ነገር በሞዴል 22 ወይም በተቋሙ ሕጋዊ የወጪ ደረሰኝ ወጪ ማድረግ አለበት፡፡
- (ሐ) በማዘዣ ወረቀት መሰረት ለውስጥ ታካሚዎች ያደላቸውን የናርኮቲክ መድኃኒትን በቅጽ NPS/08/A እና ሣይኮትሮፒክ ንጥረ ነገርን በቅጽ NPS/08/B መረጃ መያዝ አለበት፡፡
- (መ) በማዘገና ወረቀት መሰረት ለተመሳሳሽ ታካሚዎች ያደሳቸውን ናርኮቲክ መድኃኒት በቅጽ NPS/09/A እና ግይኮትሮፒክ ንጥረ ነገር በቅጽ NPS/09/B መረጃ መያዝ አለበት፡፡

4) ማንኛውም የመድኃኒት ንግድ ተቋም የናርኮቲክ ወይም የግይኮትሮፒክ ንጥረ ነገርን የገዛበትን ኢንቮይስ በፋይል እንደቅደም ተከተላቸው በመመዝገብ ከአምስት ዓመት ለማያንስ ጊዜ መረጃ መያዝ አለበት፡፡

16.የሪፖርት አቀራረብ

- 1) ማንኛውም የመድኃኒት ንግድ ተkም ፣
 - (ሀ) ያስመጣውን የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገርን በቅጽ NPS/01/A፣ NPS/01/B እንደ ቅዴም ተከተላቸው በየሩብ አመቱ መጨረሻ ለባለስልጣኑ ሪፖርት ማድረግ አለበት፡፡
 - (ለ) የሳከውን የናርኮቲክ መድኃኒትን ወይም የግይኮትሮፒክ ንጥረ ነገርን በቅጽ NPS/02/A፣ NPS/02/B እንደ ቅደም ተከተሳቸው በየሩብ አመቱ መጨረሻ ለባለስልጣት ሪፖርት ማድረግ አለበት።
 - (ሐ) ያስፋፈለውን የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገርን በቅጽ NPS/03/A፣ NPS/03/B እንደ ቅደም ተከተላቸው በየሩብ አመቱ መጨረሻ ለባለስልጣኑ ሪፖርት ማድረግ አለበት፡፡
 - (መ) የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገርን በቅጽ NPS/04/A፣ NPS/04/B እንደ ቅደም ተከተላቸው በዓመቱ መጨረሻ ለባለስልጣት ሪፖርት ማድረግ አለበት፡፡

2) ማንኛውም አምራች

- (ሀ) ያመረተውን የናርኮቲክ መድኃኒትን ወይም የግይኮትሮፒክ ንጥረ ነገርን በቅጽ NPS/01/D፣ NPS/01/E እንደ ቅዴም ተከተላቸው በየሩብ አመቱ መጨረሻ ለባለስልጣኑ ሪፖርት ማድረግ አለበት።
- (ሰ) ያስመጣዉን የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንዋሪ ነገርን በቅጽ NPS/02/D ፣ NPS/02/E እንደ ቅዴም ተከተላቸው በየሩብ አመቱ መጨረሻ ለባለስልጣኑ ሪፖርት ማድረግ አለበት፡፡
- (ሐ) የሳከዉን የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገርን በቅፅ

- NPS/02/A፣ NPS/02/B እንደ ቅዴም ተከተላቸው በየሩብ አመቱ መጨረሻ ለባለስልጣት ሪፖርት ማድረግ አለበት፡፡
- (መ) ያከፋፌለውን የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንዋሪ ነገርን በቅጽ NPS/03/A እና NPS/03/B እንደ ቅደም ተከተሳቸው በየሩብ አመቱ መጨረሻ ለባለስልጣት ሪፖርት ማድረግ አለበት፡፡
- (ሥ) የናርኮቲክ መድኃኒትን፣ ሳይኮትሮፒክ ንጥረ ነገርንና ጥሬ እቃዎች በቅጽ NPS/04/A, NPS/04/B እና NPS/03/D እንደ ቅዴም ተከተሳቸው በዓመቱ መጨረሻ ለባለስልጣት ሪፖርት ማድረግ አለበት።
- 3) ማንኛውም የጤና ተቋም ስለገዛውና ስላደለው የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገር አስመልክቶ በዓመቱ መጨረሻ በቅጽ NPS/15/A እና NPS/15/B እንደ ቅዴም ተከተላቸው ለክልል ጤና ቢሮ ሪፖርት ማድረግ አለበት::
- 4) ማንኛዉም የክልል ጤና ቢሮ በስሩ ከሚገኙ የጤና ተቋማት ስለተሰራጨ የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገር መረጃ በመሰብሰብ እስከ ጥር 30 በቅፅ NPS/16/A እና NPS/16/B ለባለስልጣኑ ሪፖርት ማድረግ አለበት፡፡
- 5) ማንኛውም አስፔሻላይዝድ የጤና ተቋም ስለገዛውና ስላደለው የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገር በዓመቱ መጨረሻ በቅፅ NPS/15/A አና NPS/15/B አንደ ቅደም ተከተላቸው ለባለስልጣኑ ወይም ለባለስልጣኑ ቅርንጫፍ ፅ/ቤት ሪፖርት ማድረግ አለበት።

ክፍል አምስት ስለ አስተዳዳራዊ እርምጃዎች

17.ፌቃድ ስለማባድና ስለመሰረዝ

1) ልዩ ፍቃዱ የሚሰረዘው፣

- ሀ) ድርጅቱ አግባብ ባለው አካል በንግድ ሥራ ወይም አገልግሎት እንዳይሰግራ መከልከሉ ሲረ*ጋ*ገጥ፣
- ለ) ድርጅቱ ልዩ ፍቃዱ ከተሰጠበት አላማ ተቃራኒ ተማባር ሲፈጸም ከተገኘ፣
- ሐ) ልዩ ፍቃድ የተሰጠባቸው መመዘኛዎች ሲጓደለ፣
- መ) በአንቀጽ 4 በተጠቀሰው መሰረት ባለፍቃዱ ተፋተኛ ሆኖ ከተገኘ፣
- 2) ፍቃድ ሲታገድም ሆነ ሲሰረዝ ባለስልጣኑ ጉዳዩ የሚመለከታቸው መ/ቤቶች እንዲያውቁት ያደርጋል፡፡
- 3) በተሰጠው የጊዜ ገደብ ሥራ ላይ ያልዋለ ልዩ ፍቃድ በቂ ምክንያት ወይም ማስረጃ ሲቀርብበትና አሳማኝ ሆኖ ሲገኝ ብቻ ሊራዝም ይችላል፡፡

18. ልዩ ፌቃድን ስለመመለስ

ማንኛወም ድርጅት፤

- 1) ልዩ ፌቃዱ ከታገደ ፣ ከተሰረዘ ወይም ሳይታደስ ከቀረ፤
- 2) በስሙ ልዩ ፌቃዱ ያወጣው ባለሙያ ከሞተ ወይም
- 3) የሚሰጠዉ አገልግሎት ለሕብረተሰቡ ጤና አደገኛ ነዉ ብሎ በባለሥልጣኑ ከታመነ ወይም ድንገተኛ የሕብረተሰብ የጤና ችግር ሊፌጠር ይችላል ተብሎ ከታመነ፤ ልዩ ፌቃዱን በሁለት ቀን ዉስጥ ለባለስልጣኑ መመለስ አለበት.

19. ስለቅሬታ አቀራረብ

ማንኛዉም ሰዉ የ**ልዩ ፌቃድ** ወረቀት አሰጣጥ እድሳት፣ እገዳና ስረዛ ወይም ሌሎች ባለሥልጣኑ የሚወስዳቸዉ እርምጃዎችን በተመለከተ ቅር ከተሰን አቤቱታውን በአንድ ወር ጊዜ ዉስጥ ባለስልጣኑ ሳደቋመዉ ቅሬታ ሰሚ አካል ማቅረብ ይችላል፡፡

20. እኅዳና ስረዛ ስለማንሳት

ማንኛውም የብቃት ማረ*ጋገጫ* ምስክር ወረቀት የታገደበት ወይም የተሰረዘበት ድርጅት የተጣለበት አገዳ ወይም ስረዛ ሲነሳለት የሚችለው በዚህ መመሪያ አንቀፅ 19 መሰረት ቅሬታ አቅርቦ ቅሬታው ተቀባይነት አግኝቶ ሲወሰን ይሆናል፡፡

21. የናርኮቲክ መድኃኒት ወይም የሣይኮትሮፒክ ንጥሬ ነገር ስለማስተሳለፍ

ማንኛውም ስራውን በራሱ ፌቃድ ያቋረጠ፣ እገዳ ወይም ስረዛ የተጣለበት ድርጅት በእጁ ያሉ አገልግሎት ላይ መዋል የሚችሉ የናርኮቲክ መድኃኒት ወይም የሣይኮትሮፒክ ንጥረ ነገር ባለስልጣኑን በማስፌቀድ ልዩ ፌቃድ ሳሳቸው ድርጅቶች በተሰጠው ጊዜ ገዴብ መሸጥ፣ ማከፋፊል ወይም ማስተሳለፍ አለበት፡፡

ክፍል ስድስት

ልዩ ልዩ ድን*ጋጌዎች*

22. ስለ አያያዝ

ማንኛውም የጤና ተkም የናርኮቲክ መድኃኒትን ወይም የሣይኮትሮፒክ ንጥረ ነገር፣ ኢንቮይስ፣ የሪኮርድ ቅጽ መዛግብት፣ የማዘዣ ወረቀትና የመሳሰሉትን ቁልፍ ባለው የብረት ግጥን ወይም በተለየ ክፍል ውስጥ ማስቀመጥና ቁልፉም በተፈቀደለት ባለሙያ ብቻ እንዲያዝ ማድረግ አለበት።

23.ስለ አወጋገድ

- 1) ማንኛውም የአገልግሎት ጊዜው በማብቃቱ ወይም በመበላሽቱ የተነሳ ተቅም ላይ የማይውል የናርኮቲክ መድኃኒት ወይም የሣይኮትሮፒክ ንጥረ ነገር ወይም ጥሬ ሪቃ አወጋገድ ስርዓት እንደሚከተለው ይሆናል፣
 - (ሀ) ተጠሪነታቸው ለክልል ጤና ቢሮ የሆኑ ጤና ተ<u>k</u>ማት ለሚገኙበት ክልል ጤና

ቢሮ ዞን ወይም ለወረዳ የጤና ጽ/ቤት ጥያቄ በማቅረብ እንዲወገድ ማድረግ አለባቸው፡፡ ስለመወገዱም በቅጽ NPS/14 የምስክር ወረቀት በመስጠት ለባለስልጣኑ ሪፖርት ማድረግ አለባቸው፡፡

- (ለ) ተጠሪነታቸው ለባለስልጣኑ ወይም ለባለስልጣኑ ቅርንጫፍ ጽ/ቤት ጥያቄ በማቅረብ እንዲወገድ ማድረግ አለባቸው፡፡ ስለመወገዱም በቅጽ NPS/14 የምስክር ወረቀት ይሰጣል፡፡
- 2) በሕገወጥ ዝውውር፣ ምርት ወይም ገበያ ውስጥ የተገኙ ወይም የተያዙ የናርኮቲክ መድታኒት ወይም የሣይኮትሮፒክ ንጥረ ነገር ዓለም አቀፍ ስምምነቶች በሚደነግጉትና በባለስልጣኑ የመድታኒት አወጋገድ መመሪያ መሰረት እንዲወገዱ ይደረጋል።

24.የተሻሩ ሀጎች

- 1) የናርኮቲክና ሳይኮትሮፒክ መድሀኒቶችን ለመቆጣጠርና በአግባቡ ጥቅም ላይ ለማዋል የወጣ መመሪያ ጥር 1996 በዚህ መመሪያ ተሽሯል፡፡
- 2) ይህንን መመሪያ የሚቃረን ማንኛዉም መመሪያ ወይም የአሥራር ልምድ በዚህ መመሪያ ውስጥ የተመለከቱ ጉዳዮችን በሚመለከት ተፈጻሚነት አይኖረዉም፡፡

25.መመሪያው የሚጸናበት ጊዜ

ይህ መመሪያ ከጥቅምት 1 ቀን 2005 ዓ.ም ጀምሮ ተፈጻሚ ይሆናል፡፡

የሁሉ ደነቀዉ ዋና ዳይሬክተር

የኢትዮጵያ የምግብ የመድኃኒት የጤና ክብካቤ አስተዳደርና ቁጥጥር ባለሥልጣን

INTRODUCTION

WHERE AS, it is found necessary to deter the illicit production, distribution and use of narcotic drugs and psychotropic substances

WHERE AS, it is found necessary to regulate the import, export, distribution, prescription, dispensing and use of narcotic drugs and psychotropic substances;

WHERE AS, to achieve these ends it is essential to lay down a secured narcotic drugs and psychotropic substances control system;

NOW THEREFORE in accordance with the Food, Medicine and Health Care Administration and Control Proclamation No. 661/2009 and Article 55(3) a Directive is hereby issued as follows.

PART ONE

GENERAL

1. Short Title

This Directive may be cited as "Directive to Control and promote proper use of

Narcotic Drugs and Psychotropic Substances No. 17 therein

2. Definitions

In this Guideline, unless the context provides otherwise;

- 1. "Narcotic Drug" shall mean any drug subject to control according to Narcotic Drugs conventions of the United Nations ratified by Ethiopia. This shall also include a drug that is categorized as narcotic drug by the Food, Medicine and Health Care Administration and Control Authority.
- "Psychotropic Substance" shall mean any substance subject to controlaccording to psychotropic substances convention of the United Nationsratified by Ethiopia. This shall also include a substance that is categorized aspsychotropic substance by the Food, Medicine and Health Care Administration and Control Authority.
- 3. "Health Institution" shall mean any governmental, non-governmental or private institution that carry out promotive, preventive, curative andrehabilitative activities or medicine trade or services;
- 4. "Authority" shall mean the Food, Medicine and Health Care Administration and Control Authority of Ethiopia.
- 5. "Appropriate organ" "appropriate organ" means, as the case may be, the executive organ or a state government organ authorized to implement food, medicine and controllable health related institution administration and control activities at a region level or other organ authorized by law.
- 6. "Drug trading institution" shall mean an institution authorized or issued certificate of competence and/or trade license by the Authority and/or otherconcerned body to produce, import, export, whole sale and retail of Narcoticdrugs, psychotropic substances or precursor chemicals.

PART TWO

LICENSE AND LICENSING PROCEDURE

3. Requirements for Licensing

- 1. No person shall manufacture, import, export, distribute, hold, or store narcotic drugs and psychotropic substances or perform chemical analysis or research on same unless it is licensed.
- 2. In accordance with sup article (1) of this Article a license is issued only for a person who have a certificate of competence from the appropriate organ to import, export, manufacture, distribut or store drugs or delivering health service.
- 3. Any person desiring to oprate activities mentiond in this Article of sub article (1) shall complete the application form in accordance article 5.

4. Prohibition from Licensing

Any person who engaged on any of the following activities cannot get a license which permit to hold narcotic drugs or psychotropic substances:

- Any offence against drug abuse, illega narcotic drugs or psychotropic substances trafficking, or enactment to commit any such offence, or forgery, the use of forged documents, assumption of false identity, or anyone convicted of planning to commit any such offence.
- 2. Where there are no adequate rooms or facilities to keep narcotic drugs or psychotropic Substances.

5. Applications for Import or Export permit

- 1. The application for obtainment of a special license under Article 3 shall be submitted in duplicate to the Authority and shall contain the following information.
 - a) Name in full, profession, and nationality of the applicant.

- b) Name of the institution and its address.
- c) The kind of trade in Narcotic Drugs and Psychotropic Substances to be thought.
- d) Name, type and strength of Narcotic Drugs or Psychotropic substances or raw materials to be employed.
- 2. The Authority shall have the power to reject an application and request amendment, reduce the quantity or weight requested, or exclude certain items from the list of requisition.

6. Issuance of Special Import or Export Permit

- 1. The Authority shall issue a special license for importation of any Narcotic Drug or Psychotropic Substance or raw materials ifit finds that
 - a) The substance is included in the National List of Drugs.
 - b) The substance is necessary to provide medical and scientific needs or other legitimate needs of the country
- 2. The Authority shall issue a special license for exportation of any locally manufactured narcotic drug or psychotropic substance when the exporting institution obtain special import authorization from the regulatory authority of the importing country.
 - 3. A special Import or Export license for Narcotic drug or psychotropic substance shall be given at most within two working days following application.

7. Import

- 1. To import Narcotic Drugs and Psychotropic Substances any person shall fulfill the following requirement :
 - a) Special import permit issued by the authority.
 - b) Narcotic Drugs and Psychotropic Substances shall import ony by air and packed separately.
 - c) Invoice only for this purpose.
- 2. Any person shall attach the following documents with the application to release from the port of entry:

- a) Registration certeficate
- b) Batch Analysis Certificate (original or copy)
- c) Packing List
- d) Bill of Loading or Airway Bill
- e) Comercial Invoice
- f) Pre-import permit certificate
- 3. The authority shall approve the attached document and given permit at the port of entry.copies of the said certificate of clearance shall be given for revenue and custom authority and the importer.

8. Distribution of Copies of the Special Import Permit

The special license that the authority given to the importer shall be prepared by four copies and then:

- 1. The original copy (copy1) of the special permit shall be issued by the Authority to the importer. The importer shall transmit this copy to the foreign exporter.
- 2. The duplicate copy (copy2) shall be forwarded by the Authority to the proper governmental authorities of the exporting country.
- 3. The triplicate copy (copy 3) shall be forwarded by the Authority to the customs Authority at the port of entry for follows up.
 - 4. The quadruplet copy (copy 4) shall be retained by Food, Medicine and Health Care Administration and Control Authority

9. Special Import or export Permit & Expiration Dat

- 1. Any special license issued to import or export Narcotic drugs or psychotropic Substances shall be valid only for ninety (90) days and to import or export once.
- 2. An import or export permit being issued by the authority shall be cancelled provided no shipment has been made and returned to the authority or in theevent that a permit is lost and proven with evidence, the authority shall issue new import or export permit in replacement to the cancelled or lost one.

PART THREE

PRESCRIPTION AND DISPENSING AGAINST PRESCRIPTION

10. Prescription and Dispensing of Narcotic Drugs or Psychotropic Substances

- 1. Any Medical practitioner who has permitted to hold Narcotic Drugs or Psychotropic Substances should only prescribed by Narcotic Drugs or Psychotropic Substances prescription papers.
- 2. Dispensing narcotic drug or psychotropic substances only by narcotic drugs or psychotropic substances prescription papers.
- 3. Any Medical personnel cannot Sell or supply narcotic drugs or psychotropic substances on presentation of a prescription, where he knows that the prescription is forged, unlawfully altered, canceled or expired.
- 4. Any Medical practitioner cannot supply, sell or help to use narcotic drugs or psychotropic substances unless they are prescribed for him by medical personnel authorized to prescribe.
- 5. Any Medical practitioner cannot prescribe narcotic drugs or psychotropic substances for himself
- 6. Any Medical practitioner cannot prescribe narcotic drugs or psychotropic substances without sufficient reason or above the standard dose, even if he has a license.

11. Medical practitioners Entitled to Issue narcotic drugs or psychotropic substances

A prescription for narcotic drug or psychotropic substance may be issued only by an individual practitioner who is authorized to prescribe narcotic drugs or psychotropic substances in which he is licensed to practice his profession.

12. Manner of Issuance of Prescription

- 1. Any prescription for narcotic drugs or psychotropic substances shall contain the following information:
 - a) Be dated as of, and the day when issued.

- b) Bear the full name, age, sex, address and card number of the patient, diagnosis (ICD code no).
- c) It shall bear name, strength, dosage form quantity of the drug; date of prescribing and clear direction for use.
- d) Bear the name, address, signature, license number and signature of the practitioner.
- e) Bear the name, address, signature of the dispensers.
- f) It shall bear the official seal of the Health Institution from which it is prescribed.
- 2. Prescription papers shall be written with ink or indelible pencil and signed by the Practitioner

13. Dispensing of Narcotic Drugs or Psychotropic Substances

- 1. Any Medical practitioner cannot supply narcotic drugs or psychotropic substances unless the prescription paper shall not fulfill requirements mentioned as Article 14 sub.article (1) of this directive.
- 2. A prescription containing narcotic drug or psychotropic substance shall not be dispensed after the elapsing of fifteen days as from the date on which it was issued.
- 3. Any Medical practitioner should not dispense or supply narcotic drugs and psychotropic substances for Children (under 15), his family or others.

PART FOUR

RECORDS AND REPORTS

14. General

- Any person, who manufactured, imported, exported or distributed narcotic drugs and psychotropic substances should keep record of his daily activity.
- 2. Any persone licensed to manufacture, import, and export or distribute narcotic drugs and psychotropic substances or raw material shall send a report about the drugs or raw materials to the authority.

15. Records

- 1. Any narcotic drugs and psychotropic substances trade institution shall kept records of :
 - a) Imported Narcotic Drugs on form NPS/01/A;
 - b) Imported Psychotropic Substances on form NPS/01/B;
 - c) Exported Narcotic Drugs on form NPS/02/A;
 - d) Exported psychotropic substances on form NPS/02/B;
 - e) Distributed Narcotic drugs on form NPS/03/A
 - f) Distributed psychotropic substances on form NPS/03/B
- 2. Any narcotic drugs and psychotropic substances manufacturing factory shall kept records of :
 - a) Imported and used Narcotic drugs and psychotropic substances raw materials on FormNPS/02/D
 - b) Manufactured Narcotic drugs on Form NPS/01/D
 - c) Manufactured Psychotropic substances on Form NPS/01/E
 - d) Distributed Narcotic drugs on form NPS/03/A
 - e) Distributed psychotropic substances on form NPS/03/B
- 3. Any health institution shall keep records of:
 - a) Purchased or donated narcotic drugs or psychotropic substances, on Model 19 or valid goods reciving memo of the inistitution.

- b) Distributed narcotic drugs or psychotropic substances to dispensary pharmacies, on model 22 or valid siv of the inistitution.
- c) Dispensed narcotic drugs to in-patients, on form NPS/08/A psychotropic substances on form NPS/08/B.
- d) Dispensed to outpatients on the grounds of a prescription, narcotic drugs on form NPS/09/A and psychotropic substances on form NPS/09/B.
- 4. Every drug trade shall keep records of purchased narcotic drugs or psychotropic Substances and all invoices related to them in a chronological file for not less than five Years.

16. Reports

- 1. Every drug trade inistititions shall report for the authority about:
 - a) Imported narcotic drugs and psychotropic substanses on form NPS/01/A and NPS/01/B respectively, at the end of every quarter:
 - b) Exported narcotic drugs and psychotropic substanses on form NPS/02/A and NPS/02/B respectively, at the end of every quarter:
 - c) Distributed narcotic drugs and psychotropic substanses on form NPS/03/A and NPS/03/B respectively, at the end of every quarter:
 - d) Raw materials of of narcotic drugs and psychotropic substanses on form NPS/04/A and NPS/04/B respectively at the end of the year.

2. Every manufacturer

- a) Manufactured narcotic drugs and psychotropic substanses on form NPS/01/D, NPS/01/E respectively report for the authority at the end of every quarter.
- b) Imported narcotic drugs and psychotropic substances on form NPS/02/D, NPS/02/E respectively report for the authority at the end of every quarter.
- c) Exported narcotic drugs and psychotropic substances on form NPS/02/A and NPS/02/B respectively report for the authority at the end of every quarter.

- d) Distributed narcotic drugs and psychotropic substance on form NPS/03/A and NPS/03/B respectively report for the authority at the end of every quarter.
- e) Narcotic drugs and psychotropic substance raw materials on form NPS/04/A, NPS/04/B and NPS/03/D respectively report for the authority at the end of the year.
- Every Health Institution shall send reports of purchased and dispensed narcotic drugs and psychotropic substances at the end of every year onForms NPS/15/A and NPS/15/B respectively to the respective regional health bureaus.
- 4. Every Regional health bureaus shall send the summary of all the reports that are collected from health institutions, under their supervision, to the Authority on form NPS/16/A and NPS/16/B until January 31.
- 5. Every specialized health institution shall send the reports of purchased and dispensed narcotic drugs and psychotropic substances on Forms NPS/15/A and NPS/15/B respectively to the authority or its branches at the end of every year.

PART FIVE

Administrative Measures

17. Suspention and revocation of license

- 1. The authority shall suspend or revoke the license on any of the following grounds
 - a) The Institution prohibited by the appropriate organ to engaged on any commercial business.
 - b) The Institution engaged on activities not allowed to do.
 - c) Fail to meet the license requirements.
 - d) In accordance of Art. 6 the licensee found guilty.
- 2. The authority will inform the concerned bodies when the lice suspended or revoked.

3. The time for special license may prolong upon sufficient and evidence based resone.

18. License return

The license shall be returned to the Authority within two (2) working day

- 1. When the special license suspended, revoked or not renewed
- 2. upon the death of the licensee
- 3. When the authority has shown any act which constitutes a threat to the public health or safty.

19. Complaint Handling

Any person whose license suspended or revoked can submit his complaints to be reviewed by the pannal established by the authority in accordance to directive No. 10/2005, with in 30 days from the time when decisen is rendered.

20. Change of decision

A dicision made to suspended or revoked the license shall be changed only when the person fill his complaints in accordance to Art. 20 and accepted.

21. Transfer of Narcotic drugs and psychotropic substances

Any person who ceases to operat as a business or the license suspended or revoked shall sale, distribute or transfer any Narcotic Drugs or Psychotropic substances and prescriptions under his possession to other licensed Institutions within a fixed period of time under the authority permission.

PART SIX

MISCELLANEOUS

22. Storage

Every Health Institution shall be stored Narcotic drugs and psychotropic substances invoices, registers, prescriptions and the like in a strong locked metal cupboard or in a special room the key to which shall at all times remain in the possession of the authorized pharmacy.

23. Disposal

- 1. Any Narcotic drugs, psychotropic substances or raw materials no longer useful, due to expiry or damage, shall be destroyed in the following manner:
 - a) Health institutions under regional health bureaus shall apply for the health bureaus or zone and disposed under their direct supervision and get disposal certificate on form NPS/14 and shall report to the authority.
 - b) Health institutions under the authority or its branch direct supervision shall apply for the authority or its branchs and disposed under their direct supervision and get disposal certificate on form NPS/14.
- 2. Narcotic Drugs or Psychotropic substances seized in illicit traffic, illicitmanufacture (Clandestine laboratory) and illicit market shall be disposed, in accordance with the International Treaty requirement, and in accordance of the Authority disposal directive.

24. Inapplicable Laws

No directive or practice shall, in so far as it is inconsistent with this directive, be applicable with respect to matters covered by this directive.

25. Effective Date

This directive shall enter into force from the date of signature by the Director General of the Authority

Yehulu Denekew General Director

Ethiopian Food Medicen & Health care Administration & Control Authority

FORM NPS/01/A

Quarterly Statistics of Imports of Narcotic Drugs	
Name of Reporting Organization	
Address: Region City/Town	
P.O. Box Tel	
These statistics relates to the quarter of the calendar year	_

					Stock	Import	Remar			
: (Narcotic Drug	Dosage Form	Strength	At the Beginning of the Quarter	Imported	Locally Purchas ed	Distribute d	at the end of the quarte r	Permit / No.	k
_										
-										
								_	_	

Remark:-Report on the following Narcoticdrugs is required quarterly

- 1. Morphine
- 2. Codiene Phosphate
- 3. Pethidine
- 4. Fentanyl
- 5. Methadone
- 6. Others is present

Quarterly Statistics of imports of				
Psychotropic substances				
Name of Reporting Organization _				
Address: Region	City/Town			
P.O. Box	Tel			
These statistics relates to the	quarter of the calendar year			
	Quantity	Ctook	Import	Г

					Stock	Import	Remar			
Ser. No.	Psychotr opic Substanc es/ Drugs	Dosage Form	Strength	At the Beginning of the Quarter	Imported	Locally Purchas ed	Distribute d	at the end of the quarte r	Permit / No.	k

Remark: - Report on the following psychotropic drugs is required quarterly

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam
- 7. Midazolam

- 8. Pentobarbitone
- 9. Phenobarbitone
- 10. Temazepam
- Other combination drugs containing controlled psychotropic substances

FORM NPS/02/A

Quarterly							
Name of							
Address:	Reg	gion _			City/Town		
Ρ.	O. Box			Tel.			
These sta	atistics r	elates	s to the	quarte	er of the calendar year		
					Quantity	Stock	export

						Quantity	Stock	export	Remar	
Ser. No.	Narcotic Drug	Dosage Form	Strength	At the Beginning of the Quarter	Exported	Manufact ured	Distribute d	at the end of the quarte r	Permit / No.	k

Remark:-Report on the following Narcoticdrugs is required quarterly

- 1. Morphine
- 2. Codiene Phosphate
- 3. Pethidine
- 4. Fentanyl
- 5. Methadone
- 6. Others is present

FORM NPS/02/B

Quarterly Statistics of Exports of	
Psychotropic substances	
Name of Reporting Organization	
Address: Region	City/Town
P.O. Box	Tel
These statistics relates to the	_ quarter of the calendar year

Ċ						ntity	Stock	Export	Remar	
No.			_	At the				at the	Permit/	k
Ser.	Psychotro	Dosage Form	Strength	Beginning	Exported	Manufact	Distributed	end of	No.	
S	pic	osage Form	rer	of the		ured		the		
	Substanc	ے ت	St	Quarter				quarter		
	es/									
	Drugs									

Remark: - Report on the following psychotropic drugs is required quarterly

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam
- 7. Midazolam

- 8. Pentobarbitone
- 9. Phenobarbitone
- 10. Temazepam
- Other combination drugs containing controlled psychotropic substances

Quarterly Statistics of distributed

	Narcotic Dru	ugs							
	Name of Re	eporting O	rganization	·					
	Address: Re	egion		Cit	y/Town				
	P.O. Bo	ох		_ Tel					
	These statis	stics relate	s to the c	alendar ye	ear(Quarter of t	the calenda	ar of the year	r
Ser. No	Narcotic	Drugs	Strength	Dosage form	Date of Issue	Issued to	Quantity Issued	Issuing/ transfer Voucher No.	Remark
	Domark: F	Poport on	the following	na Narcat	ic Drugs is re	auirod Oua	rtorly		
	Remark F	Report Off	THE TOHOW	ng maicol	ic blugs is le	quired Qua	iiterry.		

- 1. Codeine Phosphate
- 2. Morphine
- 3. Methadone

- 4. Fentanyl
- 5. Pethidine
- 6. Other Controlled

Substances if present

7. Midazolam

	Quarterly Statistics	of distribu	ıted					
	Psychotropic Subst	tances						
	Name of Reporting	g Organiza	tion			 		
	Address: Region _			City/Town				
	P.O. Box		Tel _			_		
	These statistics re	lates to th	ne calend	ar year _	Qι	uarter of th	ne calendar	of
	the year	_						
Ser. No	Psychotropic substances	Strength	Dosage form	Date of Issue	Issued to	Quantity Issued	Issuing/ transfer Voucher No.	Remark
	Remark: - Report	 on the follo	 	/chatronic	 Substance	 s is require	 ed_guarterly	
	Alprazolam		ownig i oʻ		ntobarbiton	•	ou quartoriy	
	2. Chlordiazepoxide			9. P	henobarbito	one		
	3. Clonazepam			10. T	emazepam			
	4. Diazepam			11. O	ther combir	nation drugs	containing	
	5. Medazepam					· ·	pic substance	es
	6. Oxazepam					'		

Form NPS/02/D

Annual Statistics of Psychotropic and Narcotic				
Raw Materials				
Name of Reporting Organization				
Address: Region City/Town				
P.O. Box Tel				
These statistics relates to the calendar of the year				

	. No.	Description of the raw	Type of raw materia	Unit	Balance at the beginning of the	Quantity		Stock at the	Import	Issuin	R
						Importe d	Consum ed/Issue	end of the	permit no.	g/	е
										transf	m
	Ser.	materials	pe ma)	year		d	quarter		er	ar
	0,		Ту							vouch er	k
										no.	
										110.	
l											
L											
L											

Remark: - Report on all controlled Narcotic and Psychotropic Raw Materials is required quarterly

FORM NPS/01/D

	Name of	Reportin	ng Org	ganizatio	on					
	Address:	Region			CIŲ	y/Town				
	These sta	atistics r	elates	to the	qua	arter of the c	 alendar year			
No.	Naraatia				Quantity	T		Stock at the	Import Permit	Remar k
Ser. No.	Narcotic Drug	Dosage Form	Strength	unit	At the Beginnin g of the Quarter	manufactur ed	Distributed	end of the quarte r	/ No.	
							<u> </u>	Ĺ		

Remark: -Report of the following Narcotic Drugs is required quarterly

Quarterly Statistics of Manufactured Narcotic Drugs

1. Codeine Phosphate

4. Fentanyl

2. Morphine

5. Pethidine

3. Methadone

6. Other Controlled

Quarterly Statistics of Manufactured

Psychotropic Substances

Name of Reporting Organization ______

Address: Region ______ City/Town _____

P.O. Box ______ Tel. _____

These statistics Relates to the _____ quarter of the calendar year ____

ġ					Quantity	Stock at the	Import Permit	Remar k		
Ser. No.	Psychotr opic substanc e	Dosage Form	Strength	tinu	At the Beginning of the Quarter	manufacture d	Distributed	end of the quarte r	/ No.	

Remark:-Report of the following psychotropic substances is required quarterly

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam
- 7. Midazolam

- 8. Pentobarbitone
- 9. Phenobarbitone
- 10. Temazepam
- Other combination drugs containing controlled psychotropic substances

	HEAL [*] Name	NSED AND TH INSTITU of Health I	TION nstitution:		OTIC DRUG		PD IN				
	Descri	ption of Dru	ıg		Issued						
	Chief	pharmacist:	Name		Signatu Signatur	ıre					
	FORM NPS/08/ Date Name of Health Institution: Serial No The following is an accurate record of Total quantity each used in ward Department										
Date	Please			d clearly an			Dose				
			patient								
	Ward	physician: N	lame		Signature	e	_				
	Ward	Head Nurse	: Name		Signatur	e					

Nam Seria Desa	al No cription of D	Institution: _	Quantity	 Issued						
Chie Head	f pharmacis d Nurse: Na	nt t: Name ame		ire e						
Date Nam Seria The Tota	FORM NPS/08/B Date Name of Health Institution: Serial No The following is an accurate record of Total quantity each used in ward Department Please fill the following record clearly and neatly.									
Date	Hour	Name of patient	Bed No.	Chart No.	Nurse	Dose				
Date	1100.		Bed No.	Chart No.	Nurse	Dose				
Date	1100.		Bed No.	Chart No.	Nurse	Dose				
Date	1100.		Bed No.	Chart No.	Nurse	Dose				

Record of Dispensed Narcotic Drugs in Dispensary Pharmacy of Health
Institution
Name of Health Institution
Address
Serial No

S.No	Date	Name of	Age	Sex	Address	Description of drug	Quantity dispensed	Name of prescriber	Prescription serial No
		patient							

Remark: Record on the following Psychotropic Drugs is required

- 1. Codeine Phosphate
- 2. Morphine
- 3. Methadone

- 4. Fentanyl
- 5. Pethidine
- 6. Other Controlled

Record of Dispensed Psych Institution	otropic Drugs	in Dispensary	Pharmacy	of Health
Name of Health Institution _				
Address				
Serial No				

S.No	Date	Name of patient	Age	Sex	Address	Description of drug	Quantity dispensed	Name of prescriber	Prescription serial No
		1							

Remark: Record on the following Psychotropic Drugs is required

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam
- 7. Midazolam

- 8. Pentobarbitone
- 9. Phenobarbitone
- 10. Temazepam
- Other combination drugs containing controlled psychotropic substances

Annual Statistics of Narcotic Drugs

Name of Reporting Organization ______ City/Town ______

P.O. Box _____ Tel. _____

These statistics relates to the calendar of the year ______

							Quantity	Stock	Remar
Ser. No.	Narcotic Drug	Dosage Form	Strength	At the Beginning of the Year	Imported	Locally Purcha sed	Distributed /consumpti on during the year	at the end of the year	k

Remark: -Report on all controlled the following Narcotic Drugs isrequired annually

- 1. Codeine Phosphate
- 2. Morphine
- 3. Methadone

- 4. Fentanyl
- 5. Pethidine
- 6. Other Controlled

Form	NPS/04/B

Annual Statistics of	Psychotropic											
Substances												
Name of Reporting Organization												
Address: Region		City/Town										
P.O. Box	Tel											
These statistics relates to the calendar of the year												
		O	Charl	D								

							Quantity	Stock	Remark
No.			_	At the				at the	
Ser.	Psychotr opic	Dosage Form	Strength	Beginning	Imported	Locally	Distribute	end of	
S	opic	os; For	rer	of the		Purchase	d/consu	the	
			SI	year		d	mption	year	
	es						during the year		
							trie year		

Remark:-Report on all controlled the following Psychotropic Drugs is required annually

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam
- 7. Midazolam

- 8. Pentobarbitone
 - 9. Phenobarbitone
- 10. Temazepam
- Other combination drugs containing controlled psychotropic substances

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Form NPS/03/D

Annual Statistics of Raw Materials		
Name of Reporting Organization		
Address: Region	City/Town	
P.O. Box Tel _		

				Balance at the	Quantity	0	Balance at the	Remark
Ser. No.	Narcotic or Psychotro pic raw materials	Type of raw materia	Unit	beginning of the year	Importe d	Consum ption during the year	end of the year	

Remark: -Report on all controlled Narcotic and Psychotropic substance raw materialsis required annually

Annual Report of Narcotic Drugs
Name of Reporting Health institution

FORM NPS/15/A

	P.O.	Box _			Tel								
	These statistics Relates to the calendar year												
Ser. No.	Narcotic Drug	Dosage Form	Strength	Balance at the Beginning of the Year	Quantity purchase d during the year	Purchas ed from	consumpti on during the year	balance at the end of the year	Remark				

Address: Region _____ City/Town _____

Remark: -Report on the following Psychotropic Drug is requiredAnnuallyat the end of December.

- 1. Codeine Phosphate
- 2. Morphine
- 3. Methadone

- 4. Fentanyl
- 5. Pethidine
- 6. Other Controlled

Annual Report of Psychotropic Substances

Name of Reporting Health Institution ___

FORM NPS/15/B

	Address:	Region			_City/Towr	າ							
	P.O. Box			Tel									
	These statistics Relates to the calendar year												
Ser. No.		Dosage Form	Strength	Balance at the Beginning of the Year	Quantity purchas ed during the year	Purchase d from	consumpti on during the year	balance at the end of the year	Remar k				
							-						

Remark:-Report on the following Psychotropic Drug is required annually at the end of December.

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam
- 7. Midazolam

- 8. Pentobarbitone
- 9. Phenobarbitone
- 10. Temazepam
- Other combination drugs containing controlled psychotropic substances

Annual Report of Narcotic Drugs

FORM NPS/16/A

Address: City/Town													
	P.O. Box Tel												
	These statistics Relates to the calendar year												
Ser. No.	Narcotic Drug	Dosage Form	Strength	Balance at the Beginning of the Year	d during	consumpti on during the year	balance at the end of the year	Remark					

Name of Reporting Region _____

Remark:-Report on the following Psychotropic Drug is required annually at the end of December.

- 1. Codeine Phosphate
- 2. Morphine
- 3. Methadone

- 4. Fentanyl
- 5. Pethidine
- 6. Other Controlled

FORM NPS/16/B

Annual Report of Psychotropic Substances Name of Reporting Region							
Address:City/Town							
P.O. Box Tel							
These statistics Relates to the calendar year							

Ser. No.	Psychotr opic substanc e	Dosage Form	Strength	Balance at the Beginning of the Year	Quantity purchas ed during the year	consumpti on during the year	balance at the end of the year	Remark
						_		_

Remark:-Report on the following Psychotropic Drug is required annually at the end of December.

- 1. Alprazolam
- 2. Chlordiazepoxide
- 3. Clonazepam
- 4. Diazepam
- 5. Medazepam
- 6. Oxazepam

- 7. Pentazocine
- 8. Pentobrabitone
- 9. Phenobarbitone
- 10. Temazepam
- 11. Other combination drugs containing controlled

						Ref. No Date			
	Substa We h precur	ances of the services of the s	or precurs y certify emicals e	or cher that N numera	nicals arcotic ted /imp	drug(s) oorted/	Narcotic drugs,), psychotropic stocked in rvision of insp on	substanc	ce(s) or have
S.No	discription	Unit	quantity	Batch no	Expiry date	MFD	manufacturers	Country of origin	remark
	1 2 3 Note: admin Contro Origin 2nd co		copy of to and cority				orized person Food, Medicine	——e and He	althcare