The Science and Mysteries of Sleep

Mark T Gabr, MD, FAASM
What is Sleep?

- Sleep is a reversible behavioral state of perceptual disengagement from and unresponsiveness to the environment. Sleep deprivation is typically followed by compensatory increase.

- Sleep is an active and highly organized brain state with very complex physiological and behavioral changes.

- Sleep is present in the entire animal kingdom, including insects. It has persisted throughout the evolution of mammals and birds, and is even present in lower forms.
Sleep Phylogeny

• **Reptiles, amphibians, fish** - sleep is usually determined by behavioral criteria.

• **Birds** – Sleep is very similar to that in mammals, but REM sleep is much shorter and alternates with NREM sleep in very short intervals (9 sec / 2.5 min).

• **Mammals** - all mammals sleep, and all except one species (Cetaceans) have the cyclical alternation between REM and NREM sleep.
Sleep Phylogeny

Unihemispheric sleep in the dolphin
Determinants of Sleep

- **Homeostasis**: sleep loss increases daytime sleepiness and recovery sleep.

- **Circadian Rhythms**: sleep-wake periodicity over the 24 hours of the day; controlled by the SCN.

- **Age**: the ability to sleep decreases, not the need for sleep.

- **Individual Sleep Differences**: specific value varies from person to person and may be genetically determined.
Total Sleep Requirement

Percentage of all people

Hours of sleep

- 3
- 4
- 6
- 7
- 8
- 9
- 10
- 11
What Happens in Sleep?

NREM / REM Sleep Cycle

Courtesy of R. Ristanovic, MD.

Adapted from Rogers et al. Sleep. 1994;17:590.
Electrographic State Determination

Electroencephalogram (EEG) = Brain Waves

Electrooculogram (EOG) = Eye Movements

Electromyogram (EMG) = Muscle Tension
<table>
<thead>
<tr>
<th>State</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Awake</td>
<td>Low voltage – random, fast</td>
</tr>
<tr>
<td>Drowsy</td>
<td>8 to 12 cps – alpha waves</td>
</tr>
<tr>
<td>Stage 1</td>
<td>3 to 7 cps – theta waves</td>
</tr>
<tr>
<td>Stage 2</td>
<td>12 to 14 cps – sleep spindles and K complexes</td>
</tr>
<tr>
<td>Delta Sleep</td>
<td>(stages 3 and 4) ½ to 2 cps – delta waves &gt;75 µV</td>
</tr>
<tr>
<td>REM Sleep</td>
<td>Low voltage – random, fast with sawtooth waves</td>
</tr>
</tbody>
</table>
## NREM vs. REM Sleep

<table>
<thead>
<tr>
<th>Physiologic Variable</th>
<th>NREM</th>
<th>REM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate</td>
<td>Regular</td>
<td>Irregular</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>Regular</td>
<td>Irregular</td>
</tr>
<tr>
<td>Blood pressure</td>
<td>Regular</td>
<td>Variable</td>
</tr>
<tr>
<td>Skeletal muscle tone</td>
<td>Preserved</td>
<td>Absent</td>
</tr>
<tr>
<td>Brain O2 consumption</td>
<td>Reduced</td>
<td>Increased</td>
</tr>
<tr>
<td>Response to CO2</td>
<td>Same as W</td>
<td>Depressed</td>
</tr>
<tr>
<td>Response to O2</td>
<td>Same as W</td>
<td>Same as W</td>
</tr>
<tr>
<td>Temperature</td>
<td>Homothermic</td>
<td>Poikilothermic</td>
</tr>
<tr>
<td>Penile tumescence</td>
<td>Absent</td>
<td>Present</td>
</tr>
</tbody>
</table>
REM Sleep Physiology

Tonic Components
1. EEG Desynchronization
3. EMG Atonia
4. Penile Tumesence

Phasic Components
1. Rapid Eye Movements (PGO-Spikes)
2. Muscle twitches
3. Irregular respirations
4. Cardiovascular instability
Circadian Rhythm of Sleep and Core Body Temperature
The natural endogenous circadian period of humans is slightly more than 24 hours, generally about 24.2 hours. In a time-free environment (no time cues) days are longer than 24 hours, arising one hour later every day. This is called free-running rhythm.

Keeping the basic 24-hour cycle involves daily synchronization of the internal clock with the shorter solar day, a process known as entrainment. It involves advancing the internal clock by a fixed time period (about 0.2 hour) every day. This adjustment is derived from exposure to environmental time cues (Zietgebers).

Czeisler et al, science, 1999
• The suprachiasmatic nucleus (SCN) is the central master pacemaker (Biological Clock) which regulates sleep-wake cycle and other circadian rhythms. It is a paired structure of 10K neurons each, located at the base of the anterior hypothalamus above the optic chiasm.

• Light is the strongest time cue that synchronizes circadian rhythms with the environment. Light reaches the SCN via the retinohypothalamic tract (RHT).
Melatonin & The Circadian System

Melatonin is the endocrine secretion most closely linked to the circadian system. It is actively secreted by the pineal gland for 10-12 h at night in the dark. Melatonin secretion is activated by darkness and suppressed by light. The SCN regulates the secretion of melatonin by the pineal gland, and melatonin itself has a feedback effect on the SCN by decreasing its firing rate, thus promoting sleep.
Sleep Neurophysiology
“Sleep Centers”

- **Cortex**
- **Thalamus**
  - Cortical Activation
  - Sleep Spindle
  - EEG Synchronization
- **Hypothalamus**
  - Sleep/Wake Switch
- **SCN**
  - Circadian Clock
- **Brainstem**
  - Ascending Cortical Activation
  - REM/SWS Switch

Courtesy of E. Mignot, MD. Stanford University.
# Sleep Neurochemistry

Neurotransmitters and Sleep/Wake Promoting Substances

<table>
<thead>
<tr>
<th>Wakefulness</th>
<th>NREM Sleep</th>
<th>REM Sleep</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetylcholine</td>
<td>GABA VLPO Neurons</td>
<td>Acetylcholine REM-on</td>
</tr>
<tr>
<td>Norepinephrine</td>
<td>Galanin</td>
<td>Glycine Atonia</td>
</tr>
<tr>
<td>Histamine</td>
<td></td>
<td>Monoamines (NE, HA, 5-HT) REM-off</td>
</tr>
<tr>
<td>Serotonin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypocretin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dopamine</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Why do we sleep?
Many Proposed Theories – No Answers

We know a lot about what happens if we do not sleep, but very little on the reason for sleep.

- **Restoration** - Animals with high metabolic rates sleep longer.
- **Memory Processing** - Decision making, reasoning and episodic memory improve with sleep.
- **Immune Theory** - Sleep deprivation affects the immune system.
- **Growth Theory** - SWS increases growth hormone production.
- **Energy Conservation** - Rest allows the animal to be more prepared to activity when awake.
- **Preservation** - Keeping quite when it is dark may prevent the animal from being seen by predators.
Functions of Sleep

Sleep is essential for survival

- Total sleep deprivation in rats leads to death within 10 – 20 days, faster than if they were totally deprived of food but slept normally. Fatal Familial Insomnia leads to death after several months.

- Sleep drive in humans is so strong that sleepiness increases with small reductions in nightly sleep times and sleep loss can cause severe functional impairments.

- All studied land mammals show maximal sleep at birth, leading to the conclusion that sleep is required for brain development.
Mammals sleep per 24 Hours

<table>
<thead>
<tr>
<th>Animal</th>
<th>Total Sleep</th>
<th>REM Sleep</th>
</tr>
</thead>
<tbody>
<tr>
<td>Horse</td>
<td>3.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Elephant</td>
<td>3.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Giraffe</td>
<td>4.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Seal</td>
<td>6.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Man</td>
<td>8.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Baboon</td>
<td>8.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Dolphin</td>
<td>10.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Cat</td>
<td>12.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Rat</td>
<td>12.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Opossum</td>
<td>16.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Bat</td>
<td>20.00</td>
<td>0.50</td>
</tr>
</tbody>
</table>
Functions of Sleep:
NREM Sleep: Neural Integrity

- There is huge variation in the amount of sleep that different species need, and that is not related to evolutionary classification. The opossum sleeps for 18 hours a day, whereas the elephant gets by with only 3 hours.

- Animal size is the major determinant of the amount of sleep! There is greater sleep requirement in species with low energy reserve. Small animals have higher metabolic rates and higher brain temperatures resulting in large amount of free radicals with increased cellular injury.

- NREM sleep may allow brain cells to repair membranes damaged by free radicals during waking (neuronal integrity).
Sleep Ontogeny

Functions of Sleep: REM Sleep: Neural Maturation

- The amount of REM sleep is highest early in life and in immature animals. The champion REM sleeper is the Platypus with 8 hours a day in REM sleep, while the dolphin has no REM sleep! Michel Jouvet’s theory: Genetic programing of neuronal connections (neural maturation).

- REM sleep deprivation can alleviate clinical depression, thus mimicking the effect of SSRI antidepressants. REM sleep may allow monoamine receptors (Serotonin, norepinephrine and histamine) to rest and regain full sensitivity which is crucial during waking for mood regulation.

- Contrary to popular belief, REM sleep has not been proven to play a role in memory consolidation or learning ability. People who have brain damage preventing REM sleep, or who have drug-induced blockade of REM sleep, have normal—or even improved—memory.
Disorders of Sleep

Mark T Gabr, MD, FAASM
Sleep Disorders are common

- Estimated 70 million Americans have clinically significant sleep problems
- Prevalence of insomnia about one-third of adult population
- Obstructive Sleep Apnea prevalence in patients with essential hypertension is over 25%
Sleep Disorders are Serious

• Chronic insomniacs report decreased quality of life, memory and attention problems, decreased physical health

• Increased incidence of motor vehicle crashes in patients with OSAS

• Total direct costs of insomnia in U.S. $13.9 billion (1995)
Sleep Disorders are Treatable

Multiple successful treatment modalities exist, including pharmacotherapy, surgical intervention, behavioral therapy, continuous positive airway pressure.
Sleep Disorders are Underdiagnosed

• Less than 14% of medical interns questioned patients about sleep (Haponik, 1996)

• Survey of 222 V.A. patients; no sleep symptoms recorded despite 47% prevalence (Meissner, 1998)

• Survey of 10 million ICD dx: 100,000 cases OSAS expected; 73 diagnosed (1992)
Sleep Disorders - Socioeconomic Consequences

- 40 million Americans suffer from chronic disorders of sleep.

- 95% of these remain unidentified and undiagnosed.

- The annual direct cost of sleep-related problems is $16 billion, with an additional $50-$100 billion in indirect costs (accidents, litigation, property destruction, hospitalization, and death).
Sleep Disorders - Socioeconomic Consequences

- More than 100,000 motor vehicle accidents annually are sleep-related.
- Disasters such as Chernobyl, Three Mile Island, Challenger, and Exxon Valdez were officially attributed to errors in judgment induced by sleepiness or fatigue.
Disorders of Sleep

- **Hypersomnia**: Disorders of Excessive Sleepiness

- **Insomnia**: Difficulty initiating and maintaining sleep

- **Parasomnia**: Abnormal behavior during sleep; Disorders of Arousals from NREM and REM Sleep
Excessive Daytime Sleepiness
Challenges in the Diagnosis

- Need to differentiate EDS from fatigue or tiredness
  - Sleepy patients often present with “lack of energy,” “tiredness,” or “fatigue” complaint
- Need to distinguish between EDS and the lethargy of depression
- Denial of symptoms is common
- Patients may be unaware of symptoms if onset is insidious
- Multiple causes of EDS

EDS is often unrecognized

Epworth Sleepiness Scale

How likely are you to doze off or fall asleep in the following situations, in contrast to feeling just tired? This refers to your way of life in recent times. Even if you have not done some of these things recently, try to work out how they would have affected you. Use the following scale to choose the *most appropriate number* for each situation.

<table>
<thead>
<tr>
<th>Situation</th>
<th>Chance of Dozing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sitting and reading</td>
<td></td>
</tr>
<tr>
<td>Watching TV</td>
<td></td>
</tr>
<tr>
<td>Sitting inactive in a public place (eg, a theater or a meeting)</td>
<td></td>
</tr>
<tr>
<td>As a passenger in a car for an hour without a break</td>
<td></td>
</tr>
<tr>
<td>Lying down to rest in the afternoon when circumstances permit</td>
<td></td>
</tr>
<tr>
<td>Sitting and talking to someone</td>
<td></td>
</tr>
<tr>
<td>Sitting quietly after a lunch without alcohol</td>
<td></td>
</tr>
<tr>
<td>In a car while stopped for a few minutes in traffic</td>
<td></td>
</tr>
</tbody>
</table>

0 = would *never* doze  
1 = *slight* chance of dozing  
2 = *moderate* chance of dozing  
3 = *high* chance of dozing

ESS Scores in Select Conditions

PLMD = periodic leg movement disorder; OSA = obstructive sleep apnea; IH = idiopathic hypersomnia.

Obstructive Sleep Apnea
Types of Sleep Disordered Breathing

- **Apnea**
  Cessation of airflow $\geq 10$ seconds

- **Hypopnea**
  Decreased airflow $\geq 10$ seconds
  associated with:
  - Arousal
  - Oxyhemoglobin desaturation
Measures of Sleep Apnea Frequency

- **Apnea Index**
  
  # apneas per hour of sleep

- **Apnea / Hypopnea Index (AHI)**

  # apneas + hypopneas per hour of sleep
Prevalence of Sleep Apnea

30-60 years old

Percent of Population

Prevalence in Co-Morbid Patients

- Drug-Resistant Hypertension: ~80% (Logan et al. J. Hypertension 2001)
- Type 2 Diabetes: 72% (Einhorn et al. Endocrine Pract 2007)
- Congestive Heart Failure: 80% (Maisel et al. HFSA 2007)
- Atrial Fibrillation: ~50% (Somers et al. Circulation 2004)
- All Hypertension: 35% (Sjostrom et al. Thorax 2002)
- Coronary Artery Disease: ~30% (Schafer et al. Cardiology 1999)
- Angina: ~30% (Sanner et al. Clin Cardiology 2001)
Sleep Apnea & Metabolic Syndrome

- Sleep-disordered breathing is at the center of Metabolic Syndrome
- Treating SDB may improve co-morbidities
Pathophysiology of Obstructive Sleep Apnea
Clinical Consequences

Obstructive Sleep Apnea

Sleep fragmentation, Hypoxia / Hypercapnia

Excessive daytime sleepiness

Cardiovascular Complications

Morbidity

Mortality
Clinical Consequences: Excessive Daytime Sleepiness

- Increased motor vehicle crashes
- Increased work-related accidents
- Poor job performance
- Depression
- Family discord
- Decreased quality of life
Consequences: Automobile Accidents

Accident/ Driver/ 5years

Consequences: Automobile Accidents

Risk of Traffic Accident: OSA + ETOH

Clinical Consequences: Cardiovascular

- Systemic hypertension
- Cardiac arrhythmias
- Myocardial ischemia
- Cerebrovascular disease
- Pulmonary hypertension
Cardiovascular Consequences: Hypertension

Prospective Study of Association between OSA and Hypertension

Odds Ratio

Apnea / Hypopnea Index (AHI)

0 0.1 - 4.9 5 - 14.9 > 15

Adjusted for age, sex, BMI, neck circ., cigs., ETOH, baseline HTN.

Consequences: Cardiovascular Disease

Cross Sectional Study of Association between OSA and CVD

Adjusted for age, sex, race, BMI, HTN, cigs., Chol.

Adapted from Shahar E et al. Am J Respir Crit Care Med 2001;163.
Diagnosis: History

- Snoring (loud, chronic)
- Nocturnal gasping and choking
  - Ask bed partner (witnessed apneas)
- Automobile or work related accidents
- Personality changes or cognitive problems
- Excessive daytime sleepiness
- Risk factors
Sleep Apnea Risk Factors

- Obesity
- Increasing age
- Male gender
- Anatomic abnormalities of upper airway
- Family history
- Alcohol or sedative use
- Smoking
Diagnosis: Physical Examination

- Upper body obesity / thick neck
  - ≥ 17” males
  - ≥ 16” females
- Hypertension
- Obvious airway abnormality
Behavioral Interventions

- Lose weight
- Avoid alcohol and sedatives
- Avoid sleep deprivation
- Avoid supine sleep position
- Stop smoking
Weight Loss

- Should be prescribed for all obese patients
- Can be curative but has low success rate
- Other treatment is required until optimal weight loss is achieved
Medical Interventions

- Positive airway pressure
  - Continuous positive airway pressure (CPAP)
  - Bi-level positive airway pressure
- Oral appliances
- Other (limited role)
  - Medications
  - Oxygen
Positive Airway Pressure
Surgical Alternatives

- Reconstruct upper airway
  - Uvulopalatopharyngoplasty (UPPP)
  - Laser-assisted uvulopalatopharyngoplasty (LAUP)
  - Radiofrequency tissue volume reduction
  - Genioglossal advancement
  - Nasal reconstruction
  - Tonsillectomy

- Bypass upper airway
  - Tracheostomy
Insomnia

A complaint of:

- Difficulty falling asleep
- Difficulty staying asleep
- Poor quality sleep

Associated with:

- Distress
- Impaired function
Insomnia: Daytime Complaints

- Fatigue, sluggishness
- Sleepiness
- Somatic complaints (aches & pains)
- Stress about poor sleep
- Mood disturbances
- Poor concentration
- Impaired performance
Insomnia
Pathophysiology

- Increased physiological arousal
  - Metabolic rate
  - Sympathetic activation
  - Hypothalamic-pituitary-adrenal axis

- Increased alertness
  - Conditioned state of hyperarousal
  - Decreased sleep efficiency
  - Normal MSLT
Insomnia Consequences

- Decreased quality of life
- Increased healthcare costs
- Increased absenteeism
- Decreased productivity
- Increased risk for developing psychiatric disorders
- Increased accident risk
Insomniacs Under-Report Sleep Problems to Physicians

Percent of insomniacs who discussed any sleep problems with their physicians

- Never discussed (69%)
- Visited specifically to discuss sleep problem (5%)
- Discussed sleep during visit for other purpose (26%)

Adapted from Ancoli-Israel S et al. SLEEP 1999;22.
Insomnia
Major Risk Factors

• Previous history of insomnia
• Increasing age
• Female gender
• Psychiatric symptoms and disorders
• Medical symptoms and disorders
Evolution of Insomnia

Insomnia
Etiology-Based Classification

- **Secondary insomnia**
  - Acute psychological, social stressors
  - Medical, neurological, psychiatric disorders
  - Medications, substances
  - Specific sleep disorders (e.g. sleep apnea, restless legs syndrome)

- **Primary Insomnia**
  - Psychophysiological
  - Sleep state misperception
  - Idiopathic
Psychophysiological Insomnia

Difficulty falling asleep or maintaining sleep

Behavioral and cognitive efforts to promote sleep

Physiological activation

Frustration, fear of sleeplessness
Behavioral Treatment: Sleep Hygiene

- Regular sleep / wake schedule
- Avoid stimulants and stimulating behavior
- Establish relaxing bedtime routine
- Provide conducive sleep environment
- Limit daytime naps
- Reduce or eliminate alcohol and caffeine
- Obtain regular exercise
- Avoid clock watching
Behavioral Treatment: Stimulus Control

- Use bed for sleep (and sex)
- Go to bed only when sleepy
- Get out of bed when unable to sleep
- Wake up at a consistent time (including weekends)
- Do not take daytime naps
Behavioral Treatment: Sleep Restriction

• Determine average time asleep
• Set time in bed = time asleep
• Consistent wake-up time
• No daytime naps

• If time asleep > 90% of time in bed then increase time in bed (15-30 minutes)
• If time asleep < 80% of time in bed then decrease time in bed (15-30 minutes)

Spielman AJ et al. SLEEP 1987;10.
Behavioral Treatment: Cognitive Therapy

Adapted from Morin CM. J Psychosom Res 1999;46.
Insomnia Pharmacotherapy

- Benzodiazepine
- Non-Benzodiazepine receptor agonists
- Antidepressants
- Antihistamines
- Melatonin
- Others
### Pharmacotherapy:
**Benzodiazepine Receptor Agonists**

**Benzodiazepines**
- Estazolam (Prosom)
- Flurazepam (Dalmane)
- Quazepam (Doral)
- Temazepam (Restoril)
- Triazolam (Halcion)
- Clonazepam (Klonopin)
- Lorazepam (Ativan)
- Alprazolam (Xanax)
- Diazepam (Valium)

**Non-benzodiazepines**
- Zolpidem (Ambien)
- Zaleplon (Sonata)
- Eszopiclone (Lunesta)
Insomnia Pharmacotherapy Guidelines

- Select appropriate medication
- Use lowest effective dose
- Use at bedtime (or later, if indicated)
- Duration of therapy
  - Use as needed for 2 to 4 weeks
  - Reduce dose as tolerated
  - Intermittent use suggested
  - Reassess and adjust approach
- Combine with behavioral strategies
Specific Populations

- **Sleep apnea:** consider antidepressant rather than benzodiazepine receptor agonist
- **Elderly:** low-dose, short-acting benzodiazepine receptor agonist
- **Substance abuse history:** antidepressant preferred
- **Co-morbid depression or anxiety disorder:**
  - SSRI + benzodiazepine receptor agonist
  - SSRI + trazodone
  - Nefazodone, mirtazapine
Insomnia Treatment
Comparative Efficacy:
CBT, Temazepam, and Their Combination

Adapted from Morin CM et al. JAMA 1999;281.
Restless Legs Syndrome

A neurological movement disorder characterized by an irresistible urge to move the legs accompanied by uncomfortable sensations that often occur in the evening or when at rest.

# Key Diagnostic Criteria for RLS

<table>
<thead>
<tr>
<th>Key RLS Diagnostic Criteria*</th>
<th>Supportive Features*</th>
</tr>
</thead>
<tbody>
<tr>
<td>✓ <strong>Urge to move the legs</strong>—usually accompanied or caused by uncomfortable leg sensations</td>
<td>– Sleep disturbances</td>
</tr>
<tr>
<td>✓ <strong>Temporary relief with movement</strong>—partial or total relief from discomfort by walking or stretching</td>
<td>– Involuntary leg movements</td>
</tr>
<tr>
<td>✓ <strong>Onset or worsening of symptoms at rest or inactivity</strong>, such as when lying or sitting</td>
<td>– Positive family history for RLS</td>
</tr>
<tr>
<td>✓ <strong>Worsening or onset of symptoms in the evening or at night</strong></td>
<td>– Positive response to dopaminergic therapy</td>
</tr>
</tbody>
</table>

* Diagnostic criteria developed by the International RLS (IRLS) Study Group in collaboration with the National Institutes of Health (NIH).

Restless Legs Syndrome (RLS)

- Affects approximately 10% of US adults
- Sleep disturbance is often the primary reason patients seek medical attention
- Common onset after age 40
- More prevalent in women
- Positive family history in 40-50%
A leading hypothesis is brain dopamine dysfunction. This could involve changes in:
  - Dopamine receptors and their function
  - Uptake of dopamine

There is evidence for brain iron deficiency in early-onset RLS
  - Iron is vital to the function of the brain’s dopamine systems

There is a strong genetic component in early-onset RLS, with several family studies demonstrating an autosomal-dominant hereditary pattern
Secondary RLS

It is important to rule out other underlying conditions that are associated with secondary RLS:

- Low serum ferritin levels or iron deficiency
- Consider other medications that may be contributing to RLS symptoms
- Renal failure
- Pregnancy
- Peripheral neuropathies
RLS Treatment
Non-pharmacological

- Avoid Caffeine, Nicotine, and Alcohol.

- Consider effect of medications that may enhance RLS: all antidepressants except Welbuterin, neuroleptic agents, antinausea agents, and sedating antihistamines.

- Iron replacement if serum Ferritin level less than 40 µg/L.
Dopamine Agonists are the drugs of choice in most patients with RLS (Pramipexole, Ropinirole).

Low potency opioids such as propoxyphene

Opioid Agonist such as Tramadol

Benzodiazepines

Gabapentin

High potency Opioid (Methadone) for refractory cases
Parasomnias
Dissociated States of Wakefulness and Sleep

- Wake
- REM Sleep
- NREM Sleep
REM-Sleep Behavior Disorder
(Wake-REM Dissociation)

Definition:

Intermittent absence of REM sleep atonia, allowing acting out of dreams
REM-Sleep Behavior Disorder (RBD)

Clinical Features:

- Older male predominance
- Progressive course, preceded by prolonged prodrome
- Dramatic, potentially violent, motor behavior during sleep, emerging >60-90 min after sleep onset.
- Chief complaint: injury to self or bed partner during sleep.
- Dream-enacting behavior always correlates with reported dream. Completely discordant with waking personality.
RBD - Altered Dreams

- Vivid, intense, full of action, unpleasant.
- Dreamer is being threatened or attacked by unfamiliar people, animals, insects.
- Dreamer is rarely the primary aggressor.
- Fear and anger are predominant moods.
- Dreamer is often defending his wife in a dream while actually beating her in bed.
RBD - Sleep-Related Injury

- Bruises
- Lacerations
- Fractures (including C2 “hang man”)
- Subdural hematomas
- Dislocations
- Abrasions/rug burns
- Tooth chipping, hair pulling
RBD - Prevalence

- 0.8% prevalence in the general population
- 87% of patients are men, with a mean age of onset of 52 years
- Typically a disorder of middle-aged and older males, but females and any age group can be affected
RBD - Polysomnography

REM Sleep without Atonia
Acute RBD

- Alcohol/Drug/Medication withdrawal
- Drug intoxication (Anti-cholinergic agents, Tricyclic antidepressants, MAOIs)
- Relapsing multiple sclerosis
Chronic RBD

• Idiopathic (52%)

• Associated with Neurologic Disorders (40%)

• Medication-induced: Beta-blockers, SSRIs, Venlafaxine (Effexor), Mirtazapine (Remeron), Tricyclic antidepressants– BUT NOT Bupropion (Zyban, Wellbutrin), a dopaminergic antidepressant.
RBD, Parkinsonism, and Dementia

- 45% of RBD patients develop Parkinson’s disease several years after the diagnosis with a mean of 11.5 years (range 5-23) from the onset.

- 64% of RBD patients develop Synucleinopathy: Parkinson’s disease, Multiple System Atrophy, or Lewy body Dementia.

- Men 50 years of age and older with idiopathic RBD should be told about the high risk for future Parkinsonism and Dementia.
Treatment of RBD

- Secure the environment.

- **Clonazepam** 0.5 – 1.5 mg is most effective; results in rapid control of problematic sleep behaviors and altered dreams in 90% of patients.

- Melatonin (3 – 12 mg) can be effective

- Other options: Carbamazepine, Clonidine, Gabapentin, and Donepezil.